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(54) Title: DNA SEQUENCE-BASED HLA TYPING METHOD

(57) Abstract

The present invention provides a process for determining genotypes in highly polymorphic systems by polymorphic systems by polymorphic approach of the property of the propert C). The synthesized cDNA molecules are then en-zymatically amplified using different combinations of oligonucleotides for each loans and directly se-quenced with Taq polymerase using an internal oili-gonucleotide. The sequenced genes are then analyzed.

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Technical Field

10 resulting nucleic acid ladders to determine the genotype the alleles carried by any given individual at a gene of sample nucleic acid. directly sequenced followed by evaluation of the primers. The polymerase chain reaction products are with conserved and non-conserved oligonucleotide the method of the present invention involves amplifying locus or loci of interest by polymerase chain reaction such as the major histocompatibility complex of humans, including Class I and Class II HLA genes. Specifically, determining genotypes of highly polymorphic systems, The present invention relates to a process for

Background of the Invention

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윱 25 20 antigens, 44,000 dalton polypeptides which associate of 29,000 and 34,000 daltons, respectively. These Class with B-2 microglobulin at the cell surface. target cells by helper T lymphocytes. II molecules are also involved in the recognition of cells by cytotoxic T lymphocytes. HLA Class II loci molecules are involved in the recognition of target includes the human leukocyte antigens (HLA) gene complex encode cell surface heterodimers, composed of proteins response. The various HLA Class I loci encode the HLA regulate the cell-cell interactions of the immune six. This region encodes cell-surface proteins which which is located on the short arm of human chromosome The major histocompatibility complex (MHC) The Class I

HLA-DPB and HLA-DPA loci of the HLA Class II region WHO nomenclature committee for factors of the HLA system exhibit an extremely high degree of polymorphism. The Class I region as well as the HLA-DRB, HLA-DQB, HLA-DQA, The HLA-A, HLA-B, and HLA-C loci of the HLA

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designated 25 alleles of HLA-A (HLA-A-0101, A-0201, HLA-DRB alleles, 13 HLA-DQB alleles, 8 HLA-DQA alleles, etc.), 32 alleles of HLA-B, and 11 alleles of HLA-C, 43 [Marsh and Bodmer, Immunogenetics, 31:131 (1990)]

implications of its polymorphisms (i.e., in determining its molecular basis and the functional function of the HLA molecules, much effort has gone into high degree of polymorphism is thought to relate to the 4 HLA-DPA alleles and 19 HLA-DPB alleles. Since this

10 transplantation). With the cloning of certain HLA genes this effort has extended to the DNA level.

most polymorphic genetic systems known [Bach, Immunol. short arm of human chromosome six constitute one of the The Class II genes of the HLA-D region on the

20 5 Doherty, Nature, 248:701 (1974)] or as foreign antigens 295:806 (1976)]. in alloresponses [Bach and Van Rood, N. Engl. J. Med., Today, 6:89 (1985)]. The HLA Class II molecules (DR, DQ presentation in the context of self [Zinkernagel and serve as restricting elements in nominal antigen non-covalently associated chains (alpha and beta) which and DP) are heterodimeric glycoproteins composed of two

encoded specificities can be determined by serological Allelic polymorphism of the HLA-D region

- 8 25 al., supra (1988); Erlich and Bugawan, in PCR RFLP and oligotyping [Bidwell, supra (1988); Tiercy et Tiercy et al., Proc. Natl. Acad. Sci. USA, 85:198 development of molecular approaches to typing, such as polymorphisms and, more recently, oligotyping [Bach, determination of restriction fragment length methods for phenotyping, mixed lymphocyte cultures using (1988)]. Present efforts focus largely on the supra (1985); Bidwell, Immunol. Today, 9:18 (1988); homozygous typing cells, primed lymphocyte testing,
- 35 Techniques, H. A. Erlich, ed., Stockton Press, New Yor)

polymorphisms are located in hypervariable regions of their N-terminal domains, encoded by the second exon of DQ, and -DP alieles has revealed that their amino acid The cloning and sequencing of several HLA-DR,

- ហ design of allele-specific oligonucleotides which can be DRB1, DRB3/4/5, DQA1 and DQB1, DPA1 and DPB1 genes used in the characterization of the known HLA Class II [Marsh and Bodmer, supra (1990); Todd et al., Nature, 329:599 (1987)]. This information has allowed the
- 5 a solid support (oligomer typing) or for sequencing polymorphisms by means of their hybridization to DNA on (1989); Todd et al., supra (1987); Saiki et al., Science, 230:1350 (1985); Mullis and Paloona, Methods [Tierry et al., supra (1988); Erlich and Bugawan, supra,
- 20 5 rapid, requires the use of a rather large number of Gyllenstein and Erlich, Proc. Natl. Acad. Sci. USA, Enzymol., 155:335 (1987); Saiki et al., Nature, 324:163 85:7652 (1988)]. Oligonucleotide typing, although (1986); Scharf et al., Science, 233:1076 (1986);
- oligonucleotides for each locus and cannot detect practical for the analysis of Class I polymorphisms. approach may not be easily applicable to and may not be to exist in non-Caucasian populations; further, the previously unidentified sequence polymorphisms, likely
- 25 Direct sequencing of single-stranded DNA generated by successfully used to examine polymorphism at DQA1 locus this approach to DRB genes is, however, problematic due PCR using allele-specific oligonucleotides has been [Gyllenstein and Erlich, supra (1988)]. Application of
- 30 to the strong sequence homology among DRB1, DRB3, DRB4 and DRB5 genes and the presence of up to four different versions of each of these genes in most individuals generated by direct sequencing make this present process (isotypic complexity). The very complex ladders
- မ္မ impractical for accurate and rapid determination of HLA been limited to previous knowledge of the HLA types types. Thus, direct sequencing of HLA-PCR products has

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suitable for routine HLA typing (Bach, supra (1985); Erlich and Bugawan, supra (1989)]. Bidwell, supra (1988); Fiercy et al., supra (1988); carried by a given individual and as such is

- transplantation. Rejection of organ grafts is believed recipient are identical. The numerous alleles of HLA to be diminished if the HLA alleles of donor and connection with many medical procedures, e.g., organ Currently, HLA typing is routinely done in
- ឋ 5 typing are the availability of standard sera necessary and Class II HLA loci. Other drawbacks to current HLA the polymorphisms associated with the alleles atlass I techniques are incapable of differentiating among all of genes in the population also make HLA typing useful for paternity testing. However, the currently available
- 20 typing in organ transplants and in relatively high not be highly accurate. techniques causes unnecessary delay and the results may the length of time associated with current HLA typing volume genetic evaluations, such as paternity testing, test results (i.e., MLC takes 5-7 days), and that only are detected by these techniques. In the case of tissue the already known HLA types, but not new polymorphisms, to conduct serological tests, the speed of obtaining
- 8 25 to have previous knowledge of his or her HLA types as genes carried by any given individual without the need capable of determining the nucleotide sequences of the avoids the use of oligonucleotides specific for each defined by other methods. Furthermore, the invention in the case of the HLA gene complex, a system that is the limitations imposed by previous methods. That is, eystems, such as the HLA gene complex, that addresses determine genomic information in highly polymorphic Accordingly, there is a need for a method to
- 35 requires the use of only a small number of known allele. The technique we present is rapid, oligonucleotide primers, and can readily detect new

Class I and Class III genes and is automatable. applicability to the analysis of Class II as well as sequence variants unidentifiable with more conventional approaches. This system is exemplified by its

Summary of the Invention

15 6 of the present invention involves sequence-based typing (SBT) which provides for unequivocal determination of oligonucleotide primers. conserved and non-conserved (non-allele-specific) sequencing genemic or complementary DNA molecules for each allele at each gene locus to be sequenced using polymorphic genes of a subject by amplifying and direct determining the nucleic acid sequence of one or more The present invention relates to a method for In a broad sense, the method

20 more genetic loci of interest, regardless of the of a unique locus, as exemplified by DQA or the like; for example: (1) simple homozygosity or heterozygosity complexity of the polymorphism at these loci, including, be employed to determine genetic polymorphism at one or DNA or expressed (RNA) copies of such a locus. SBT can direct, simultaneous, sequence analysis of both genomic genetic polymorphism at any genetic locus of interest by

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and C loci genes.

method is envisioned as being useful to determine A, B,

25 and closely linked copies of a locus, as exemplified by polymorphisms are of the first, and simplest, type. I genes or the like. Most known human genetic locus compounded by interlocus complexity, such as Class DRB or the like; and (3) intra-allelic complexity at a (2) isotypic complexity due to multiple, closely related

30 each copy of that locus by equal amplification and locus and direct interpretation of the overlapping direct sequencing of mixtures of both alleles of that selection of a given locus with equal representation of of interest as is exemplified by each of the types of HLA loci. The SBT strategy is designed to ensure sequence data comprised of only the copies of the locus Use of the SBT method provides overlapping

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at one or more genetic loci of interest which can be providing a method for determining genetic polymorphism employed, for example, in HLA typing, detection, sequencing patterns generated by this approach.

such as, for example, sickle cell anemia, cystic evaluation, and/or characterization of genetic diseases genetic loci associated with various cancers such as evaluation, and/or characterization of polymorphism in fibrosis, Thalassemia, and the like, and detection,

10 p53, Ras, myc, associated with carcinomas, leukemias, sarcomas or the like. Use of the method according to the present

ĸ complex class genotype of a subject in a sample (e.g., Class I or Class II). Most particularly, the method is and DPB1 genes. In the case of Class I genotypes, the locus including DRB1, DRB3, DRB4, DRB5, DQB1, DQA1, DPA1 directed to determining at least one HLA Class II gene and accurate determination of a major histocompatibility invention is exemplified by a system providing for rapid

polymorphism with the method of the present invention, nucleic acid (RNA or DNA) from a sample is isolated. To determine a gene locus nucleic acid sequence

30 25 sample nucleic acid sequence is determined by: gene locus. According to the present invention, the the case of RNA, cDNA molecules for each aliele of at anneals to a conserved region of each allele of each least one gene locus to be sequenced are synthesized by employing a locus-specific oligonuclectide primer that

all of the alleles for each gene locus and chromosome to for each allele of each gene locus to be sequenced, with polymerase chain reaction to generate sufficient product amplifying the cDNA molecules or genomic DNA by

딿 alleles of each gene locus and chromosome to be oligonuclectide primer pair, and at least one of the be sequenced being amplified with at least one conserved

sequencing (clean); sequencing directly the products of conserved primer; preparing the products of each PCR for conserved oligonucleutide primer and at least one sequenced being amplified with at least one non-

10 product for each locus and primer combination(s) to polymerase and a conserved primer specific for each allele at each gene locus of each chromosome, with an each polymerase chain reaction product to detect each locus that is sequenced; and analyzing each sequenced enzyme appropriate for DNA sequencing, such as Tag

determine the genotype of the subject.

15 chain reaction product. The analysis is conducted by by analyzing each nucleic acid single and/or overlapping gene locus sequence to known sequences for each locus, comparing the nucleotide sequence of each allele of each ladder generated for each directly sequenced polymerase product for each allele of each gene locus is determined invention the sequence of each polymerase chain reaction In a preferred embodiment of the present

20 of each allele of the gene locus amplified with a oligonucleotide primer pair to the nucleotide sequence followed by comparing the sequence of each gene locus amplified with the non-conserved/conserved

25 nucleic acid ladders for sequenced alleles can be conducted visually or using computer software. conserved oligonucleotide primer pair. Comparison of

ä Automation of the process includes isolating the sample nucleic acid with an RNA/DNA extractor; amplifying the by polymerase chain reaction using a thermocycler to synthesized cDNA molecule or the isolated DNA molecule determinations, including diagnosis of genetic diseases invention is automated for use in rapid genotype In a preferred embodiment, the process of the

ដូ sequencing the polymerase chain reaction products in an sequenced polymerase chain reaction product with the automated sequencing apparatus; and analyzing each generate the polymerase chain reaction products;

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chain reaction product sequence for each allele substraction algorithm for comparing the polymerase information and the capacity to conduct the appropriate computer having a database with allelic sequence

G pair. with a non-conserved/conserved oligonuclectide primer amplified with a conserved oligonuclectide primer pair to the nucleic acid sequence of each allele sequenced

ĸ 6 of each chromosome that is amplified. Useful single groups of oligonucleotide primers useful in the steps of cDNA synthesis, cDNA/genomic DNA amplification by Table 1 herein. strand DNA oligonucleotide primers are described in nucleotide sequence of each of the alleles at each locus polymerase chain reaction and direct sequencing of the polymerase chain reaction products to determine the The invention further relates to specific

Brief Description of the Drawings

DRB5), DQA1, DQB1, DPA1 and DPB1 genes. Sequencing experiments for DRB (DRB1, DRB3, DRB4 and Figure 1B shows a schematic of the primer Figure 1A shows a schematic of the cDNA/PCR/

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30 25 primers. used for PCR; and blank boxes represent sequencing conserved (or Type 1) primers, used for PCR; checked transcripts. Stippled boxes represent primers used binding sites on DRB, DQA1, DQB1, DPA1 and DPB1 boxed represent non-conserved (or Type 2) primers, also the cDNA synthesis reactions; black boxes represent 5

used for genomic DNA samples are shown in the Figure. germline configuration. Only those primers exclusively binding sites on DQB1, DRB, DPB1 and DPA1 genes in their Figure 1C shows a schematic of the primer

performed in a different test tube. The reactions are for peripheral blood samples. Each reaction is Figure 2A shows a flow-chart of the procedure

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named with capital letter in parenthesis; these letters correspond to those shown in Table II (combinations of primers/reaction). Only the "routine" combinations of primers are shown in this Figure.

forensic samples, where DNA is usually the only available genetic material to work with. DNA in these situations is usually isolated from hair, sperm, blood stains, etc. The combinations of primers per reaction 10 shown in the Figure correspond to the "routine" combinations only.

Figure 3 shows direct sequencing of Class II
HLA dsDNA generated using conserved oligonucleotides.
Lanes are read from left to right as G-A-T-C. 1, DOB1

Lanes are read from left to right as G-A-T-C. 1, DQB1

15 ladder for a DQB1*0201/DQB1*0302 heterozygote; 2, DQA1

ladder for a DQA1*0103 homozygous cell line; 3, DRB

ladders for a DRB1*0301, DRB3*0101/DRB1*0401, DRB4*0101

heterozygote. Positions where there is more than one
band are indicated on the side of the ladder and the

20 templates they correspond to are indicated at the top of the Figure. To read unambiguously the last 50-60 base pairs of the ladder it is necessary to electrophorese the sequencing gel for an additional hour. Note that the ladders corresponding to the genes at DRB3 or DRB4

the genes at DRB1 locus, possibly due to their lower levels of expression. These differences in intensity are generally reproducible and help read the complex overlapping patterns. The positions of the first exon base pair and codon (in parenthesis) that can be read in this Figure are indicated at the bottom of each ladder.

Figure 4 shows direct sequencing of Class II
HLA DRBI dsDNA generated using non-conserved
oligonucleotides. Lanes are read from left to winter

oligonuclectides. Lanes are read from left to right as 35 G-A-T-C. Lane 1, DRB1*0101/DRB1*1501, DRB5*0101 heterozygote cDNA amplified with primer DRB17 (selects DRB5*0101 cDNA) (left) and DRB16 (selects DRB1*0101

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cDNA) (right); lane 2, DRB1*1405, DRB4*0101/DRB1*0301, DRB3*0101 heterozygote cDNAs amplified with the 5' primers DRB17 (selects DRB1*0301 and DRB3*0101 cDNAs) (left) and DRB16 (selects DRB1*0405 and DRB4*0101)

5 (right). Positions where there is more than one band or where the two ladders generated with each primer differ are indicated on the side of the ladders.

Detailed Description of the Invention

segment of DNA, composed of a transcribed region and a regulatory sequence that makes possible a transcription. The term "gene locus" refers to the specific place on the chromosome where a gene is located. The term 15 "allele" refers to the multiple forms of a gene that can exist at a single gene locus at a single chromosome and are distinguishable from the other possible alleles by their differing effects on phenotype (detectable outward

manifestations of a specific genotype). "Haplotype"

20 refers to the specific allele composition of the genes
at multiple loci on the same chromosome. As used herein
the term "genotype" refers to the specific allelic
composition of a gene at multiple linked loci at each
chromosome (2 haplotypes).

refers to a molecule having two or more deoxyribonucleotides or ribonucleotides, preferably more than three deoxyribonucleotides. The exact number of nucleotides in the molecule will depend on the function of the specific oligonucleotide molecule. As used herein the

o specific oligonucleotide molecule. As used herein the term "primer" refers to a single stranded DNA oligonucleotide sequence, preferably produced synthetically which is capable of acting as a point of initiation for synthesis of a primer extension product which is complementary to a nucleic acid strand to be

copied or a point of initiation for sequencing a DNA molecule. In the case of primers intended for use in

prime the synthesis of extension products in the length and sequence of the primer must be sufficient to molecules by polymerase chain reaction products, the synthesizing cDNA or amplifying cDNA or genomic DNA

more preferably from about 5-20 nucleotides. Specific complexity of required DNA or RNA target templates, as length and sequence of the primer will depend on length of the primer is from about 5-50 nucleotides, presence of a polymerization enzyme. Preferably, the

15 10 important considerations. well as conditions of primer employment such as sequencing primers on the DNA template are also of base pairs separating the amplification and When nested primers are used for sequencing, the number temperature, ionic strength, and MgCl2 concentration.

conservation (i.e. less than 1-2 nucleotide variations). primer" (Type 1) refers to an oligonucleotide molecule that corresponds to a region of high DNA sequence As used herein, "conserved oligonucleotide

20 Functionally, conserved primers are capable of equally While the conserved primer need not correspond exactly one mismatch with the target nucleotide template. conserved primer will have minimal, preferably less than to the nucleotide template to which it anneals, the

25 priming the target nucleotide (cDNA, PCR product, etc.) an intended number of mismatches with the possible used herein, "non-conserved oligonucleotide primer" at high stringency conditions. In contrast to this, as (Type 2) refers to an oligonucleotide molecule that has

မှ target nucleotide sequences. The intended number of number of alleles at a given locus or at a group of characterized by their selective binding to a limited mismatches being about 1-12. Non-conserved primers are mismatches can vary with a preferred number of

μ highly homologous loci. The non-conserved primer will bind to the more complementary allele or group of alieles (two or less than two) (i.e., fewer number of

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accurate results with minimal expenditure of time and used herein are specifically designed to obtain highly primers and the number of reactions per locus or loci The specific combinations of conserved and non-conserved mismatches between primer and target template sequence).

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ŭ as sickle cell anemia, cystic fibrosis, or the like, as genes related to different human genetic disorders, such individual, such as, for example, the human HLA system, polymorphic gene systems carried by any given for determining the sequences of the alleles of The present invention is directed to a process

well as gene systems associated with various cancers,

15 exemplified by its utility for determining polymorphism such as p53, myc, or the like. The present invention is the gene cDNA molecules using a limited number of using enzymatic amplification and direct sequencing of at HLA loci, particularly Class II and Class I genes, the most polymorphic human genetic loci known today,

20 method is particularly well suited to determining primers and avoiding the use of allele specific allelic sequences of Class II HLA genes, thereby " oligonuclectides as much as possible. The present providing complete HLA Class II genotype information for

25 a subject. Using the method of the present invention complete Class II HLA typing (DR, DQ and DP) can be performed in about 16 to 24 hours or less.

Generally, the method of the present invention

80 case of RNA, generation of cDNA; cDNA or genomic DNA products; and analysis of the direct sequence and direct sequencing the cDNA amplification products is information. Generation of cDNA, emplifying the cDNA amplification; direct sequencing of amplification involves: extraction of sample nucleic acid; in the

딿 accomplished using oligonucleotide primers with specific characteristics, such as those described herein

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· Oligonucleotide Primers

The oligonucleotide primers of the present invention can be synthesized using any known suitable method, such as phosphotriester and phosphodisster methods. Narang et al., Methods Enzymol., 68:90 (1979); Brown et al., Methods Enzymol., 68:109 (1979).

Oligonucleotides can be prepared using a modified solid support such as a Biosearch 8750 DNA synthesizer.

Useful primers can also be isolated from a biological source using appropriate restriction endonucleases which cut double stranded DNA at or near a nucleotide sequence of interest for use as a primer.

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B. Extraction of Sample Nucleic Acid

In the process of the present invention any source of nucleic acid can be used as the sample nucleic acid, as long as the sample contains the nucleic acid sequence of interest. For example, the sample chosen for the present method can be RNA, DNA or a DNA/RNA 20 hybrid. Typical samples include peripheral blood

O hybrid. Typical samples include peripheral blood mononuclear cells, (PRMNC's), lymphoblastoid cell lines (LCI's), hair cells or the like. For determining human HLA Class II and Class I gene polymorphisms LCI's or PBMNC's are preferred. The nucleic acid to be isolated

25 (e.g. RNA or DNA) will depend on the source of genetic material (blood stain, hair, or peripheral blood cells). However, in the case of HLA Class II genes including DNB1-5, DQB1, DQA1, DPB1 the preferred isolated nucleic acid is total cellular RNA when the typing is to

30 be done for transplantation purposes or paternity testing. For forensic uses, genomic DNA may be the preferred genetic material in which case different primer considerations would be used. Cytoplasmic and poly(A) + RNA can also be used. It is envisioned that 35 isolation of sample nucleic acid for the present process can be automated using a DNA/RNA extractor (such as

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Model 341 DNA extractor available from Applied Biosystems, Inc.; Roster City, CA).

C. Generation of cDNA

Complementary DNA (cDNA) of the sample nucleic acid is generated using specific oligonucleotide primers and cloned reverse transcriptase following general conditions suggested by the enzyme manufacturer (Bethesda Research Laboratories, Gaithersburg, MD).

10 Specific differences in type and amount of primers used, dNTP concentrations and elongation times will be readily apparent to those of skill in the art based on the Examples that follow.

D. Polymerase Chain Reaction

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Amplification of cDNA or genomic DNA for each gene locus of interest is accomplished using the polymerase chain reaction (PCR) as generally described in U.S. Pat. Nos. 4,683,195 and 4,683,202 to Mullis.

20 The PCR consists of many repetitions of a cycle which consists of: (a) a denaturation step, which melts both strands of a DNA molecule; (b) an annealing step, which is aimed at allowing the primers to anneal specifically to the melted strands of the DNA molecule; and (c) an

25 extension step, which incorporates to the primers deoxyribonucleotides complementary to those of the strand of DNA to which the primers are annealed. The PCR process, as indicated in the Examples, can be conducted using a Thermocycler (Perkin-Elmer, Cetus, 30 Emeryville, CA).

The present invention introduces the use of non-conserved oligonucleotides in the PCR procedure specifically designed to solve the problems associated with, for example, detecting, evaluating, and/or

35 characterizing polymorphism at a polymorphic gene locus or loci of an individual. In the case of HLA typing, the use of non-conserved oligonucleotides addresses the

- designed for the amplification of specific genes with the use of oligonucleotides specific for the particular gene to be amplified. However, even using completely matched primers, in most cases the PCR is not absolutely matched primers, in most cases the PCR is not absolutely matched primers, the use of conserved primers in PCR will generate complex mixtures of templates, which upon direct sequencing will be seen as overlapping sequencing ladders, cumbersome to interpret. Therefore, genes for
- 15 which the exact nucleotide sequence information is unknown can not be achieved with an adequate level of certainty. Use of non-conserved oligonucleotides which can selectively anneal under high stringency conditions to two or fewer alleles of a gene locus or group of homologous loci can provide sequence information for the
- O homologous loci can provide sequence information for the different genes at highly homologous loci in complex haterozygote combinations. Thus, the present invention provides a method useful for determining the genotype for polymorphic gene loci. This is of particular
- 25 importance to HLA typing, and is applicable to Class I HLA typing as well as Class II typing.

The difference between non-conserved primers and allele-specific oligonucleotides resides in that the latter can only be used when the presence of a

- 30 particular allele is known, and also requires the use of a specific primer for each of the alleles of the polymorphic system. Thus, combining use of a non-conserved primer and conserved primers to amplify the separate alleles of highly homologous polymorphic gone
- 35 loci can provide simpler DNA polymerase chain reaction product combinations sufficient to allow unambiguous interpretation of direct sequencing ladders of each

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allele for genotype determinations with moderate expenditure of time and economical cost.

The conditions used for the PCR reactions are preferably the same except for the temperature used in the annualing step, which is different depending on the type of primer used, conserved (Type 1) or non-conserved (Type 2). Reactions that use the former primer type are preferably performed at 37°C in the annualing step of the cycle, whereas this step is preferably performed at 10 about 55°C to 60°C in reactions that use the later type of primers. The concentrations of primers, and buffers used will be apparent from and include the process

E. Direct Sequencing Of PCR Products

parameters described in the Examples that follow.

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Direct sequencing of double-stranded DNA generated by the PCR is accomplished using an enzyme appropriate to DNA sequencing, such as Tag polymerase, or the like, and specific combinations of reagents at appropriate concentrations. The sequencing procedure can be conducted in an automatic sequencing apparatus such as the 373A Model DNA Sequencer from Applied Biosystems Inc. (Foster City, CA). The reagents, including sequencing primers and nucleic acid termination mixtures will be understood by those of skill in the art based on the direct sequencing

Analysis Of Direct Sequenced PCR Products

procedure specified in the following Examples

30 The nucleic acid ladders resulting from direct sequencing the CDNA or genomic DNA for each gene locus of interest can be assessed visually from autoradiograms or by employing a computer programmed with nucleotides sequence information for all alleles of all haplotypes and procedures for comparing sequenced alleles and known alleles of gene loci of interest. In a preferred

embodiment of the present invention, the evaluation of

conserved primer products) with a library of known chain reaction product (i.e., conserved and noncomparison of the gene sequences of each polymerase gene locus alleles involvés a two step process: (a)

- 5 U of an allele of a gene locus amplified with a conserved information for the polymerase chain reaction product alleles; followed by (b) comparison of direct sequence other than sequencing [Marsh and Bodmer, Immunogenetics, 31:131 (1990)] as well as sequences of individual homologous cell lines very well characterized by methods genotype information such as the information obtained on
- 5 This comparison employs a substitution algorithm or information for each allels of a gene locus. visual cancellation of duplicative sequence ladder amplified with a conserved/non-conserved primer pair information to generate the specific sequence

reaction product of alleles of a gene locus or loci oligonuclectide primer pair and polymerase chain

- 8 present invention can be used to amplify and sequence HLA typing, and the like). The present process is HLA typing, including Class I, Class II, and Class III genetic disease-related genes, cancer-related genes, and known and unknown highly polymorphic systems (e.g., It is envisioned that the process of the
- 25 believed to be useful for paternity testing and forensic blot-detection systems. While in the latter only a hybridization pattern is observed, direct sequencing of length polymorphism (RFLP), DNA fingerprinting or dot medicine, with more accuracy than restriction fragment
- 8 amplified products shows the exact nucleotide sequence of the amplified genes, and hence is more accurate and
- ω speed and especially improving its accuracy. As polymorphism enalysis of DRB1, DRB3, DRB4, DRB5, DQB1 evidenced by the following Examples, sequence Class II HLA typing, reducing its costs, increasing its The method is particularly well suited for

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oligonucleotides. The approach described herein is using a limited number of conserved and non-conserved any subject of unknown HLA type by means of enzymatic amplification and direct sequencing of Class II genes DQA1, DPA1 and DPB1 genes can be rapidly performed in

- ()1 sequences or sequence microheterogeneity at the the advantage of detecting the presence of new allelic methods using oligonuclectide probes and dot blots, has technology and, as opposed to previously described entirely automatable using currently available
- 15 10 allogeneic transplantation. The method of the present Class II HLA polymorphism in studies of human disease different Class II HLA loci on graft survival after and may be of interest in the search for new Class II invention allows rapid and precise sequence analysis of analyses of the effects of sequence alielism at invention is envisioned to be useful for detailed population level. The methodology of present
- 20 intended to limit the invention. illustration in the following Examples which are not The present invention is further described by

sequence variants in large populations of subjects.

EXAMPLE I

5 Preparation of Oligodeoxyribonucl Sequence Primer Combinations Usef Sequence Primer Combinations Useful for CDNA/PCR/Sequencing Reactions of Class II HLA Genes ectide Primers and

30 belows described herewithin were synthesized as described All of the oligodeoxyribonuclectide primers

condensed to a nucleoside derivatized controlled pore Milligen-Biosearch (Novato, CA), were sequentially primers: The b-cyanoethylphosphoamidites, obtained from Automated Synthesis of oligodeoxyribonucleotide

띯 glass support using a Biosearch 8750 DNA synthesizer. Condensation cycles included detritylation with condensation with benzotriazole and capping with acetic dichloroacetic acid in dichloromethane, followed by

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methodology for oligodeoxyribonucleotide synthesis is described in Caruthers, et al., Methods Enzymol., measuring dimethoxytrityle alcohol release. minutes. Yields at each step were >99% as determined by pyridine, with each cycle time being approximately 9 anhydride and 1-methylimidazole in tetrahydrofuran and The

oligodeoxyribonucleotide primers: Deprotection and Deprotection and purification of

154:287 (1987).

15 5 at room temperature for about one hour. The solution purification of oligodeoxyribonuclectide primers was oligodeoxyribonuclectide was brought to 65°C for 16 subjected to chromatography on a C18 reverse-phase hours. Ammonia was removed and the residue was support by exposure to concentrated ammonium hydroxide oligodeoxyribonuclectide was removed from the solid al., Nucl. Acids Res., 15:397 (1987). Briefly, the containing the partially deprotected performed using the procedure described by Schulhof et

until dry, resuspended in water and quantitated by acid. The detritylated oligodeoxyribonucleotide was oligodeoxyribonucleotide by treatment with 70% acetic column (RP 304, BioRad, Richmond, VA) using a linear measuring its absorbance at 260nm. recovered by precipitation in ether, vacuum centrifuged group was removed from the HPLC-purified gradient of 14 to 20% acetonitrile in 0.1 molar ammonium/triethylamine, pH 7.0. The dimethoxytrityle

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oligonuclectide primers corresponding to specified were synthesized (see Table I below) and extensively regions of HLA Class II DQA, DQB, DRB, DPB and DPA loci Using the above procedure, the following

cDNA/PCR/Sequencing Reactions

gednetice						
Listing (Seq.)	•				
No.	Type I		Annes 1	Locus(1)	<u>Template</u>	Step
1	DQB7	5'-GGTGGTTGAGGGCCTCTGTCC-3'	105-111	DQB1	RNA	RT/PCR
2	DRB20	5'-GTGCTGCAGGGGCTGGGTCTT-3'	105-111	DRB1/3/4/5	RNA	RT/PCR
3	DQA9	5'-GGTGAGGTTACTGATCTTGAAG-3'	148-155	DQA1	RNA	RT/PCR
4	DQB13	5'-AGAGACTCTCCCGAGGATTTC-3'	1-7	DQB1	RNA	PCR/SEQ
5	DRB22	5'-CTGGCTTTGGCTGGGGACACC-3'	-4-3	DRB1/3/4/5	RNA/DNA	PCR/SEQ
б	DRB11	5'-TGTTCTCCAGCATGGTGTGTC-3'	-33/-26	DRB1/3/4/5	RNA	FCR
7	DQA10	5'-CTGTCCTCCGTGATGAGCCC-3'	-10/-4	DQA1	RNA	PCR
8	DQB932	5'-TCGCCTCTGCAGGGTCGCGCG-3'	88-94	DQB1	DNA	PCR
9	DQB931	5'-TTTAAGGGCATGTGCTACTTC-3'	11-17	DQB1	DNA	PCR .
10	DQB30	5'-ATGGGGAGATGGTCACTGTGG-3'	97-104	DQB1	RNA	SEQ
11	DRB30	5'-AGGATACACAGTCACCTTAGG-3'	97-103	DRB1/3/4/5	RNA	SEQ
12	DQB5	5'-GTAGTTGTGTCTGCACAC-3'	78-83	DQB1	RNA/DNA	SEQ
1.3	DRB12	5'-GCCGCTGCACTGTGAAGCTC-3'	87-94	DRB1/3/4/5	RNA	BEQ
14	DQA29	5'-CACGGTTCCGGTAGCAGCGGTAG-3'	82-89	DQA1	RNA	SEQ
15	DQA30	5'-TACGGTCCCTCTGGCCAG-3'	19-24	DQA1	RNA	SEQ
16	DRB1.400	5'-GCGCTTCGACAGCGACGTGG-3'	38-45	DRB1/3/4/5	RNA/DNA	SEQ
17	DRB1401	5'-GAGGTGACTCTGTATCCTGAC-3'	98-104	DRB1*0701-2	RNA/DNA	PCR
16	DRB1402	5'-GATCAGGCCTGTGGACACCAC-3'	142-148	DRB1/3/4/5	RNA/DNA	PCR
19	DRB1403	5'-CCGGAACCACCTGACTTCAAT-3'	127-133	DRB1/3/4/5	RNA/DNA	SEQ
20	DRB1406	5'-GCCAAGAGTGGGCCTCGCAGC-3'	bpl8-38- intron 3	DRB1/3/4/5	DNA	PCR
21	DRB825	5'-AACCCCGTAGTTGTGTCTGCA-3'	79-85	DRB1/3/4/5	DNA	SEQ
22	DRB824	5'-GGGGACACCCGACCACGTTTC-3'	1-7	DRB1/3/4/5	DNA	PCR
23	DPB10	5'-CGGACAGTGGCTCTGACGGCG-3'	-19/-13	DPB1	RNA	PCR
24	DPB11	5'-GTTGTGGTGCTGCAAGGGCCG-3'	105-111	DPB1	RNA	RT/PCR

Sequence Listing No.	(Seg.) <u>Type 2</u>		Anneal	Locus(i)	Template	Step
42	DR1123	5'-TTGTTGGAGCAGGATAAGTA-3'	7-13	DRB1	RNA/DNA	PCR
43	DRB24	5'-CCACGTTTCTTGGAGTACTCT-3'	5-11	DRB1	RNA/DNA	PCE
44	DRB25	5'-TTTCTTGGAGCAGGTTAAACA-3'	6-13	DRB1	RNA/DNA	PCR
45	DRB16	5'-AGATGCATCTATAACGAAGAG-3'	29-35	DRB1/3/4/5	RNA/DNA	PCR
46	DRB17	5'-AGATACTTCCATAACCAGGAG-3'	29-35	DRB1/1/4/5	RNA/DNA	PCR
47	*DQB6	5'-CTGAGCACCCCAGTGGCTGAG-3'	-8/-2	DQB1	RNA	PCR
48	*D0814	5'-CTGAGGTGCTCAGTGGCTGAG-3'	-8/-2	DQB1	RNA	PCR
40	ADADIE	C1 0#040040000000000040 31	# 7 9	non:	20114	600

Anneal

97-103

-12/-5 bp-42/-62

intron 2

bp 39/59

intron 3

bp 1-21

intron 3

bp -6/-26

introp 2 104-110

-17/-23

bp -69 to

bp 55-71

intron 3

76-82

59-65

68-74

222-228 214-220

-50 intron 2

88-94

-3/-9

Locus(1)

DPB1

DPB1

OPBI

DPB1

DPB1

DPB1

DPA1

DPA1

DPA1 DPA1

DPA1

DPAI

DPA1

DPA1

DPA1

Template

RNA

RNA

DNA

DNA

DNA

DNA

RNA

RNA

RNA

RNA

DHA

DNA

RNA

RNA

RHA

RNA

DNA/RNA

Step

SEQ

PCR

PCR

SEQ

SEQ

PCR

SEQ

SEO

PCR

PCR

SEQ

PCR

RT SEQ

SEQ

RT/PCR

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SEQ/RT

Sequence Listing (Seq.) No.

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Type 1

DPB12

DPB13

DPB14

DPB15

DPB16

DPB17

DPA14

DPA15

DPA16

DPA17

DPA10

DPA11

DPA12

DPA18

DPA19

DPA20

DPA21

5'-CTTGGAGGGGGAAACATTCAC-3'

5'-TACTGATGGTGCTGCTCACAT-3'
5'-AGAGGGAGAAACAGGATTAGA-3'

5'-GCCCTGGGGACGGGCCGGGG-3'

5'-CGGCCCAAAGCCCTGACTCAC-3'

5'-CGCTCATGTCCGCCCCCTCCC-3'

-GTCAATGTGGCAGATGAGGGT-3

5'-CATATCAGAGCTGTGATCTTG-3'

5'-CTTGGGAAACACGGTCACCTC-3'

5'-CTGCTGAGTCTCCGAGGAGCT-3'

5'-CTCTAGCTTTGACCACTTGC-3'

5'-AGTCTGAGGGTGGCAGAGAGG-3'

5'-GGCCTGAGTGTGGTTGGAACG-3'

5'-CTGGCTAACATTGCTATATTG-3'

5'-GGTCCCCTGGGCCCGGGGGTC-3'

5'-GCCAGAACGCAGAGACTTTAT-3'

5'-AACTTGAATACCTTGATCCAG-3'

All the above Type 1 primers are annealed at 37°C and the Type 2 primers are annealed at 55°C. When the latter anneal at 37°C in the PCR, they do not distinguish among allelic transcripts differing by few base pairs. This list of primers includes primers which are only used in certain situations, such as to confirm homozygosity at a particular locus whenever not expected according to the typings performed at the other linked loci. The alternative combinations of primers used in each step are described in Table II below. [(*) These primers anneal to a polymorphic region of DQBI cDNAs (codons -8 to -2) encoding the 3' end of the signal peptide which has specific nucleotide nucleotide sequences for different DQBI alleles (DQB6-DQB1*0501 and DQB1*0504-, DQB14-DQB1*0501-, DQB15-DQB1*0301-).] RT = Reverse transcription; SEQ = sequencing.

Combinations of Primers for cDRA/PCR/Sequence

oligonucleotide primers for each reaction and for each There are specific combinations of

10 U combinations. In addition, Table II includes a list of necessary sequence information for obtaining highly the "routine" combinations for a particular locus not direct sequencing, which are designed to provide all the locus, including cDNA synthesis, PCR amplification and locus which may be used to confirm results obtained with "alternative" combinations of oligonuclectides for each combinations are listed in Table II below as "routine" accurate, fast and inexpensive typing results. These

15 maps. These "unexpected" results are usually indicative can be confirmed with the use of the alternative of the existence of new alleles and/or haplotypes, which these combinations of oligonucleotides is characterized combinations of oligonucleotides. In any case, each of expected according to, for instance, known haplotypic

20 by its ability to generate an end-product (sequencing appropriate software. the naked eye or processed by computer operated under ladder) which is suitable of being accurately read by

30 25 such as in transplantation, the method uses RNA isolated the only available template. Although for each template material; for forensic purposes, however, DNA is often (RNA or DNA) different combinations of oligonucleotides from peripheral blood mononuclear cells as starting For typing purposes in the clinical setting,

are used (see Table II), the general strategy for primers for "routine" RNA and DNA analysis, essentially the same. typing, including the interpretation of the results is The specific combinations of

general overview of the HLA typing strategy is shown in Figures 1 and 2 and discussed further in Examples 2 and respectively, are described below in more detail. The

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TABLE II

Alternative E. 1 DR Ah. 2 DR AH. 2 DR AC. 2 DR AD. 1 DR	Routine S. 1 DRB12/DF T. 2 U. 2 V. 2 V. 2 V. 1 X. 1	2. DNA	TYPE CI ROUTING DR DRB30/DRB12 B. 2 DRB C. 2 DRB C. 2 DRB C. 2 DRB C. 1 DP H. 1 DP
DRB12 DRB1401 DRB1406 DRB1406 DRB1406 DRB1406	DRB1406 DRB1400***** DRB1406 DRB1406 DRB1406 DRB1406 DRB1406 DRB14 DPB14 DPB14	PCR1	Type CDMA Line DRB20 1 DRB20 2 DRB20 2 DRB20 2 DRB20 1 DQA9 1 DPA14 1 DPA14 1 DPA14 1 DPA19 1 DPA19 1 DPA19 1 DPA19 1 DPA16 1 DRB20 2 DRB20 2 DRB20 2 DRB20 3 DRB20 5 DRB20 1 DRB20 5 DRB20 1 DRB20 1 DRB20 2 DRB20 2 DRB20 1 DRB20 2 DRB20 3 DRB20 1 DRB20 1 DRB20 2 DRB20 1 DRB20 1 DRB20 2 DRB20 1 DRB20 1 DRB20 2 DRB20 1 DRB20
DRB824 DRB1402 DRB16 DRB17 DPB16	DRB22 DRB24 DRB25 DRB25 DRB23 DQB931 DQB931 DPB15 DPB15	PCR2	PCR DR611 DR623 DR625 DQ613 DQ613 DQ7410 DP415 DP415 DP418 DR816 DR827 DR816 DR816 DR817 DR817 DR817 DR817 DR817 DR817
37°C 55°C 57°C	37°C	A. T.	335555533555 333335555 3 P. C.C.C.C.C.C.C.C.C.C.C.C.C.C.C.C.C.C.
DRB825 DRB1403# DRB825/DRB12 DRB825/DRB12 DPB17	DRB12/DRB1400 DRB12/DRB1400 DRB12/DRB1400 DRB12/DRB1400 DQB5: DPB16/DPB17 DPB16/DPB17	Seq	*C DRB30/DRB12 *C DRB30/DRB12 *C DRB30/DQB5 *C DRB30/DQB5 *C DQB30/DQB5 *C DQB30/DQA29*** *C DPB12/DPB13 *C DPB20/DPA21**** *C DPB30/DRB12 *C DRB30/DRB12 *C DRB30/DRB12 *C DRB30/DRB12 *C DRB30/DQB5 *C DQB30/DQB5

³ For sequencing DRB and DQB two alternative sequencing primers are indicated, both sequencing the positive strand of DNA.

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- (**) Primer DRB22 is used to sequence the negative strand whenever new allelic sequences are identified.
- (***) Bach DQA1 sequencing primer anneals to a different strand. Reaction I uses an alternative amplification primer (pRB 22 instead of DRB11) in hypothetical situations where homozygosity may not be expected according to the rest of the haplotype. Reaction K is used for sequencing the negative strand of DQB1 in situations where new allelic sequences are identified.
- (****) Sequencing of the third exon is necessary to distinguish among certain DPA1 alleles.
- (*****) Primer DRB1400 may be used in sequencing amplified DRB genes from genomic DNA to read the sequences immediately following the 3' emplification primer: Reaction Q, R, Z and Ab are alternative combinations for confirming homozygosity at the corresponding loci which may not be expected according to the rest of the Class II haplotype.
- (*) This primer combination is used to distinguish between DRB1*0701 and DRB1*0702, which differ by a single base pair in their third exons.

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EXAMPLE II Protocol: HLA Class II "Typing" by Direct Sequencing of DEB, DOB, DOA, DPA and DPB Genes

Cell Lines and Subjects

each of the known Class II haplotypes defined at the 10th International Histocompatibility Workshop [Dupont, Hum. Immunol., 26, 3 (1989)] were provided by Dr. Miriam Segall (University of Hinnesota). Forty unrelated subjects who had been previously serologically typed for Class I and Class II antigens were also studied. The serological types of each of the subjects under study were not known to the investigator performing the sequence analysis at the time the analysis was

- affected (insulin-dependent diabetes and autoimmune thyroid disease) individuals. The sequenced haplotypes, many in heterozygote combinations, included: DR7 (n=3), DRw17 (n=26), DR4 (n=16), DRw11 (n=8), DRw8 (n=4), DR1 (n=6), DRw15 (n=6), DRw16 (n=2), DRw14 (n=2), DR212 (n=3), DR5x6 (n=3). The cell lines and heterozygote combinations tested are shown in Table III. Since the complexity at DPA and DPB loci is similar to that of DQ genes, the primer combinations for DPA and
- 25 DPB typing were optimized in a smaller group of homozygote and heterozygote subjects.

HLA-DRB, DOB and DOA Transcript Amplification Using Conserved and Non-Conserved Oligonuclectides

from 5-10x10⁶ peripheral blood mononuclear cells (PBKNC) or lymphoblastoid cell lines (ICLs) by cesium chloride centrifugation [Chirgwin et al. <u>Biochemistry</u>, <u>18</u>, 5249 (1979)]. Alternatively, total RNA from peripheral blood 55 (2-10 ml) was partially purified using a much faster protocol [Gouth, <u>Anal. Biochem.</u>, <u>173</u>, 93 (1988)]. One microgram of total cellular RNA was reverse transcribed with Moloney leukemia virus reverse transcriptase

period. A 5'-primer (20 pmols) (Type 1 or Type 2 MgCl2, 0.1% gelatin) were added after the incubation buffer (500 mM KCl, 100 mM Tris-Cl, pH 8.3, 7.5-15 mM volume for 30-45 min at 37°C. Eight µl of 10% PCR specific non-sense primer (Table II) in a 20 ml final units, Promega), 75 HM each dNMP and 10 pmols of a the presence of the ribonuclease inhibitor RNAs in (10 Tris HCl, pH 8.3, 75 mM KCl, 10 mM DTT, 3mM MgCl2, in (MLVRT) (200 u, Bethesda Research Laboratories) in 50 mN

5 5 or 55°C and 30 sec at 72°C using a Perkin-Elmer Cetus Scharf et al., <u>supra</u> (1986)]. The primers used here, Faloona, <u>supra</u> (1987); Saiki et al., <u>supra</u> (1986); Thermocycler [see Saiki et al., supra (1985); Hullis and subjected to 35 cycles of 30 sec at 94°C, 30 sec at 37°C μ l with distilled water. The reaction mixture was were also added and the final volume was adjusted to 100 primars, respectively, see Table II) plus 10 pmols of the non-sense primer and two units of Taq polymerase

25 20 they anneal are shown in Table II. The reactions for same tube. DPB) can be successfully performed simultaneously in the each locus are usually performed in separate microfuge and PCR reactions for all loci (DRB, DQA, DQB, DPA and tubes. However, when using conserved primers, the cDNA their corresponding sequences and the regions to which

Direct Sequencing of Amplified Products with Tag

မ္မ µl of 1X Tag sequencing buffer (50 mH Tris~HCl, pH 9, 10 retentate (20 μ l) was dried down and resuspended in 15 unincorporated dwrps and excess of oligonuclectides by 100 (millipore) microconcentrators. One half of the spin-dialysis using Centricon-100 (Amicon) or Ultrafree-The reaction mixture (100 µl) was freed of

ü mM HgCl2). Internal oligonucleotides were used for genes, respectively (Table II). Primers for sequencing priming the sequencing of DQB, DRB, DQA, DPB and DPA each strand are listed in Table II. Only one strand is

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end-labelled with 10 pmol of gamma-P32 labelled ATP sequence is suspected. Eighty to 100 ng of primer were strand is performed in cases where a new allelic routinely sequenced for typing; sequencing of the other

labelled ATP, boiled for 5 min., and then left at room sequencing mixture without extraction of unincorporated volume. Ten ng of primer (1 μ l) were added to the polynucleotide kinase (Fromega Biotec) in a 10 μl final (5000 Ci/mmol, 10 µCI/µL) and 5 units of T4

15 5 ddGTP; b) Term mix ddA: 15 microM each dGTP, dATP, 4 µl of each of the stop nucleotide mixes: a) Term mix the annealed primer/template mixture were later added to temperature for 15 min. Eight units of recombinant Taq polymerase (USB) were added to the mixture. Four $\mu 1$ of 15 microM each dGTP, dATP, dCTP, dTTP; 45 microM

20 two consecutive periods of 10 min. at 72-74°C. After a 7.5 µM mixture of ATP, GTP, TTP, CTP, and allowed to was stopped by adding 4 ml of 95% (vol/vol) formamide/20 proceed for 5 min. After spinning down, the reaction the second cycle, each reaction was chased with 2 µl of microM ddCTP. The reactions were allowed to proceed for each dGTP, dATP, dCTP, dTTP; 1200 microX ddCTP; d) Term mix ddC: 15 microM each dGTP, dATP, dCTP, dTTP; 450 dCTP, dTTP; 600 microM ddATP; c) Term mix ddT: 15 microM

8 25 mM EDTA, beated to 80°C for 5 min. and loaded on a .4 mm min, dried, and exposed to KodaK X-Omat film for 4 to 12 (vol/vol) glacial acetic acid/5% (v/v) methanol for 15 was performed at 2500 V for 2 hr, the gel fixed in 5% thick 6% polyacrylamide/7M urea gel. Electrophoresis

Sequence-Based Typing of DR and DQ Polymorphic Genes in Homozygous Typing Cells

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initial test of the methodology. In total, these cell Histocompatibility Workshop (Table III) were used as an from the panel of the 10th International Homozygous lymphoblastoid cell lines (LCLs)

lines were representative of most of the known DR and DQ alleles at the time the analysis was conducted.

Total cellular RNA isolated from homozygous LCLs was reverse-transcribed and the resultant cDNAs amplified using conserved oligonucleotides specific for DRB1/DRB3/DRB4/DRB5 or DQB1 or DQA1 genes as described in the preceding protocol. The conserved or Type 1 oligonucleotide primers anneal to regions of conserved DNA sequences; these regions are identical among the

10 known alleles at each locus and flank the second exon of Class II genes. These conserved primers, as opposed to non-conserved or Type 2 primers, are designed to amplify all known alleles at DRB, DQA1 and DQB1 loc1 and, thus, all possible combinations of these alleles in any given

15 heterozygote. The Type 1 oligonuclectides did not cross-amplify templates at loci other than those specified by the oligonuclectides (i.e., the DQA1 primers did not amplify DRB or DQB1 transcripts and vice versa); as expected, the DRB primers also amplified any

20 DRB3, DRB4 or DRB5 transcripts present in addition to DRB1. Sequencing of these amplified templates was performed using a Type 1 primer annealing to a conserved region of the cDNAs internal to the sequence recognized by the amplification primers. Figure 1A shows the

25 general strategy for the method (SBT) and Figure 1B shows the relative position of each of the oligonucleotides used for the cDNA, PCR and sequencing reactions on the mature DNB, DQA and DQB mRNA molecules. The sequences of these primers, the loci they are

30 specific for, the specific positions (codons) to which they anneal and the reaction(s) they are used in are indicated in Table II where the specific combinations of primers that can be used for the CDNA/PCR/sequencing reactions for each locus are identified. As noted in

35 the legend to Table II, some of the primer combinations shown represent alternatives which may be useful in confirming results for a particular locus which do not

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fit with the expected sequences usually found with the rest of the haplotype. Each CDNA/PCR reaction is usually performed in a separate tube. However, when using Type I primers, the cDNA/PCR reactions for all the loci (DRB, DQA, DQB, DPA and DPB) can be performed

simultaneously in the same tube. The products of each locus are sequenced in separate tubes. Following the conditions described in the above protocol, the sequence ladders between the sequencing primer and the 5' amplification primer could be clearly read starting from

2 to 14 bases from the sequencing primer binding site.

No anomalous amplification products or sequencing ladders were detectable upon direct sequence analysis of amplified DRB, DQB1 and DQA1 cDNAs from the 43

15 homozygous cell lines tested (Table IIIa). The specific

15 homozygous cell lines tested (Table IIIa). The specific alleles at each Class II HIA locus composing the haplotypes carried by each of these cell lines are shown in Table IIIb. The number of ladders generated for each cell line was always that expected according to the

20 specificity of the amplification primers (one DQBI and one DQAI ladder for all cell lines, one DRB ladder for DRI and DRw8 cell lines and two DRB ladders for haplotypes of the DRw52 and DRw53 supertypic groups).

Thus, analysis of the homozygous typing cell lines showed that the Type I primers used for cDNA synthesis,

sequencing reactions allowed for accurate amplification and sequencing of all the tested alleles at each of these loci.

Cell I	Lines and Heteroxygote		Combinations Tested	
	58 II		- 1	
Line		Subject	ĸ	
SA	DR1-DW1	S1	DR1-Dw1/DRw17	
MZ070782	DR1-Dw20	S2		
KAS011	DRw16-Dw21	S3		
*CALOGERO	DRw16-Dw-	S4	DRw15-Dw2/DRw17	
*WJR076	DRw16-Dw21	85	DRw15-Dw2/DR4-	
*DEM	DRW16_Dw21/ne4	n h	1	
WT24	DRw16-Dw21	57	DRSY68/DBW17	
REF	DRw16-Dw22	in (*DBv13_	
Dw18/DRw17		٠,	7.6	
	DRw15-Dw2	68	DRw13-Dw19/DRw17	
8 LA	DRw15-Dw2	S10	4/DR	
*AMAI	DRw15-Dw2	S 11	Day 4	
E4181324	DRw15-Dw12	S12	DR4-Dw13/DR1-Dw1	
MT14B	DR4-Dw14	1 E	DR4-Dw13/DRw17	
Dw2		,		
RSH	DRw18-DwRSH	S15	*DR4-Dw15/DRw17	
DEC	DR4-Dw4	S16	DRW11-DW5/DRW17	
E TU	DR4-Dw4	S17	DRw12/DR1-Dw1	
JEAR	DR4-Dw13	S18	DRw12/DRw8.1	
7 AN	DR4-DW10	819	. 25	
SPOOLO	DRW11-DB2	22.0	DR4-DW4/DR7	
JBUSH	DRW11-Dw5	S22	DRw8.1/DR7	
TISI	DRw11-DwTISI	S23		
KAL	DRW11-DWJVK	S24	٧,	
BM16	DRw12-DB6	S25	٣.	
*H0301	DRw13-Dw19	826	DRw8.3/DRw15-Dw2	
NT47	DRw13-Dw19	175	DRY/DRI-DWI	
TEM	DRw14-Dw9			
EX	DRw14-Dw9			
WIVE	DRw14-Dw16			
2 F				
CP96	DR7-Dw7			
BER	DR7-Dw7			
BBC	DR7-Dw11			
250	DAWO-DWO. I			
TILA	DR R DW C C C C C C C C C			
TAB089	DRw8-Dw8.3			
DKB				

The allelic composition at DRB, DQA1 and DQB1 loci for the sequenced haplotypes corresponded to that expected according to published sequence information from well

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characterized homozygous cell lines unless indicated (*).

- Haplotypes carrying new allelic sequences (DRB1, DRB3, DQA1 or DQB1 loci).
- Only the tested heterozygote combinations are listed. The remainder of the 40 subjects tested were homozygotes or carried the haplotypes listed in this table.
- This DRB specificity (DR5x6) has been given this arbitrary designation according to serological, RFLP and sequence information.

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Table IIIb

Mail 9001 40101 40501	Kaplotype	WS#	DRB1	DRB3	DRB4	DRB5	D081	1700
00 9002 21 9012,9007 15 1501	⊢	9001	1010	•	•	•	*0501	*0101
221 9009,-64,-15 *1501 - *0201 *0502 22 9015,9007 *1501 - *0202 *0301 23 9010, 017 *1501 - *0202 *0301 24 9010, 017 *1501 - *0202 *0301 25 9010, 017 *1501 - *0101 - *0101 25 9010, 017 *1501 - *0101 - *0101 25 9025 *0401 *0201 - *0101 - *0201 26 9025 *0402 - *0101 - *0201 - *0201 27 9026 *0402 - *0101 - *0201 - *0202 28 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9036 *0402 - *0101 - *0202 29 9037 *0404 *0201 - *0202 29 9057 9054 *1402 *0201 - *0101 - *0202 29 9057 *0406 *0701 - *0101 - *0202 29 9057 *0406 *0701 - *0101 - *0202 29 9057 *0406 *0701 - *0101 - *0202 29 9057 *0406 *0701 - *0101 - *0202 29 9057 *0406 *0402 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *0101 - *0202 29 9057 *0406 *0400 - *0101 - *	0	9002	2	•	ı	ı	250	*0101
222 9012,9007 *1601 - *0202 22 9013,9017 *1501 - *0202 23 9018 *1501 - *0101 31 9085 *10301 *0101 - *0101 9025 *0401 *0401 - *0101 - *0101 9029 *0401 *0401 - *0101 - *0101 9028 *0404 *0401 *0101 - *0101 9028 *0404 *0401 *0101 *0101 *0101 9028 *0404 *0101 *0101 *0101 *0101 9028 *0404 *0101 *010		-64,-1	160	•	ı	*0201	0502	*0102
22 9013,9017 *1501	_	9007	160	ı	1	*0202	*0503	****
22 9013,9017 *1501			*1602	ì	•	*0207	*0.01	10201
#22 9010 #23 9011 #1502 #1501 #1502		,9017	*1501	1	•	*0101	+0602	0103
3 9081			*1501	•	•	1	+0602	*0102
\$\frac{3}{3}\$ \text{908}\$ \text{*1301} \text{*201} \text{*201} \text{*201} \text{*201} \qu			*1502	•	•	*D102	*0601	*0103
SEXT 3085 **0301 **0201 - **0407 **0407 **04001 **04001 **0301 **0300 **04001 **0300 **04001 **0300 **0300 **04001 **0300 **0300 **04001 **0300 **0300 **04002	3		*0301		•	1	*0201	*1501
RESH 9021 **3302 **0101 **0401 **0401 **0401 **0401 **0401 **0401 **0401 **0401 **0401 **0401 **0401 **0401 **0401 **0401 **0400 **0401 **04000 **04000 **04000 **04000 **04000 **04000 **04000 **04000 **04000 **04000 **04000 **	CASAG		1060*	0	1	t	*020I	+0501
9025 **0401 **0101 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0301 **0302 **0302 **0402 **0101 **0302 **0302 **0402 **0101 **0302 **0302 **0402 **0101 **0302 **03	DVRSH		*0302	0	ı	•	*0402	*0401
9029 90401 - *0101 - *0301 9029 9039 9039 9039 9039 9039 9039 9039			*0401		1010+	•	*0301	*0301
9030 9030 -0403 -0101 -0300 9026 9026 -0402/6 -01011 -0300 9028 9028 90402 -04001 -04000 9028 9028 90402 -04001 -04000 9028 90405 -04005 -04001 -04000 9028 9035 91102 9020 -04000 -04000 9039 91102 9020 -04000 -04000 9039 91102 9020 -04000 -04000 9039 9120 9039 9120 9039 9120 9039 9120 9030 -04000 -04000 -04000 9030 9030 9030 9030 9030 9030 9030			*0401	1	*0101	1	*0302	+030I
9024 +0403/6 - +0101 +0306 9028 +040404 - +0101 - +0306 9028 +040404 - +0101 - +0306 9028 +040404 - +0101 - +0306 9028 +040405 - +0101 - +0206 9036 +1104 +0201 - +0201 - +0206 9039 +1101 +0201 - +0306 9042 +1103 +0201 - +0306 9042 +1103 +0201 - +0306 9043 +1201 +0201 - +0306 9045 +1202 +0201 - +0306 9045 +1202 +0201 - +0306 9046 +1202 +0301 - +0406 9047 +1202 +0301 - +0406 9047 +1202 +0301 - +0406 9047 +1202 +0301 - +0406 9047 +1202 +0301 - +0406 9047 +1202 +0301 - +0406 9047 +1202 +0301 - +0406 9047 +1202 +0406 - +0406 9047 +1202 +0406 - +0406 9047 +1202 +0406 - +0406 9047 +1202 +0406 - +0406 9047 +1202 +0406 - +0406 9047 +1202 +0406 - +0406 9047 +1202 +0406 9047 +1202 +0406 - +0406 9047 +1202	·w		·	•	+0101	•	*0301	*0301
9028	DKT2	9024	0403/	1	1010	l	*0302	*0301
## 100	DWIO	9026	*0402	1	1010	11	*0302	±0301
NT3 (a) +0405 - +0101 - +0405 2 9035 +1104 +0201 - +0500 2 9035 +1101 +0201 - +0500 3 9032 +1102 +0201 - +0301 5x6 +0303 +1201 +0201 - +0301 5x6 +0303 +1201 +0201 - +0301 5x6 +0301 +1302 +0201 - +0301 5x6 +0301 +0101 - +0301 5x6 +0301 +0101 - +0502 5x6 +1301 +0101 - +0502 5x6 +1302 +0301 - +0502 5x6 +0301 +0301 - +0502 5x6 +0301 - +0301 - +0502 5x6 +0301 - +0301 - +0502 5x7 +0302 - +0303 - +0303 5x7 +0302 - +0303 5x7 +0303	Dw14	9028	*0404	t	1010	ı	+0302	±0301
2 9036	DW15 KT3	(<u>a</u>)	*0405	•	10101	1	*0401	*0301
9035 1101 1020 - 0301 1030 1030 1030 1030 1030 1030 1		8	*0405	1	*0101	1	*0201	*0301
1 9042			1100	10701	ı	,	¥0502	*0102
TWE 9039 **1102 **1201 **0301 **5x6 **1302 **0301 **0301 **5x6 **1302 **0301 **0301 **03001 **1302 **0301 **04001 **03001 **1301 **0101 **05001 **03001 **1301 **0101 **05001 **03001 **1301 **0101 **05001 **03001 **1301 **0101 **05001 **03001 **1301 **0101 **05001 **03001 **03001 **0101 **0201 **0201 **03001 **03001 **03001 **0201 **0201 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001 **03001		9042	1103	*0201	٠,		10201	1000
5 9038 *1201 *0201 - *0301 *02	MACAG		*1102	*0201		1	*0301	*0501
5x6 DR5x6 DR5x6 18 9055 18 9055 18 9062 18 9063 18 9063 18 9063 18 9063 18 9063 18 9063 18 9063 18 9063 18 9063 18 9063 18 9063 18 9063 18 9064 18 9064 18 9064 18 9064 18 9064 18 9064 18 9064 18 9065 18 9064 18 9065 18 9066 18	12-DB6		*1201	*0201	•	ı	*0301	*0501
18 9062 +1302 +0301 - +0605 18 (b) 9063 +1301 +0101 - +0502 18 (b) 9063 +1301 +0101 - +0502 19 9063 +1302 +0301 - +0502 19 9064 +1302 +0301 - +0503 10 9064 +1401 +0201 - +0503 10 9064 +0701 - +0101 - +0201 10 9096,9046 +0701 - +0101 - +0201 10 9096,9046 +0701 - +0101 - +0201 10 9096,9046 +0701 - +0101 - +0201 10 9096,9046 +0701 - +0101 - +0201 10 9096,9046 +0701 - +0101 - +0201 10 9096,9046 +0701 - +0101 - +0201 10 9096 +0701 - +0101 - +0201 10 9096 +0701 - +0101 - +0201 11 9067 +0803 +0402 12 9070 +0803 +0402 13 9076 +0803 +0301 14 9066 +0901 - +0101 - +0301 15 9066 +0901 - +0101 - +0301 16 9068 +0901 - +0301				±0101	1	,	*0301	*0501
18 9062 *1301 *0101 - *0502 19 9063 *1302 *0301 - *0603 19 9063 *1302 *0301 - *0603 19 9057,9054 *1401 *0201 - *0503 10 9064 *1401 *0201 - *0301 *0301 10 9064 *1402 *0101 - *0301 *0301 10 9096,9046 *0701 - *0101 *0301 10 9096,9046 *0701 - *0101 *0301 10 9097 *0701 - *0101 *0101 *0301 10 9050 *0701 - *0101 *0101 *0301 10 9050 *0701 - *0101 *0402 10 9050 *0803 - *0402 10 9066 *0803 - *0402 10 9066 *0803 - *0402 10 9066 *0803 - *0402 10 9066 *0803 - *0402 10 9066 *0803 - *0402 10 9066 *0803 - *0403	-Dw19			*0301	1	•	D086.5	*0102
18 (b) +1301 +0101 - +0504 + 1301 +0101 - +0504 + 1301 +0201 - +0504 + 1401 +0201 - +0503 + 1401 +0201 - +0503 + 1401 +0201 - +0301 + +0301 + +0301 + +0301 + +0301 + +0301 + +0301 + +0303 + +0301 + +0303 + +0301 + +0303 +	Dw18	062		1010*	1	,	*0603	£0103
9 9057,9054	STAG	_	1301	*0101	ı	1	+0502	*0102
Y Y Y Y Y Y Y Y Y Y	DW19	63	1302	1000	1	•	*0604	*0102
16 9064 1407 10101 10201	640	57,905		+0201	1	١	*0503	DOA1.4
9096,9046 *0701 *0101 *0				*0201	•	1	DQB5.4	S)
9094,9093 *0701 *10101	4+PA-10	2	*1402	1010	1	,	10301	1050+
9052 *0701 *01014 *0101		900	10701	•	1010-	,	1020	1020
9950 +0701 +0101 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		3	10701		:	•	1000	1010
1 9967 +0801	Dw17	9050	+0701	•			1000	1
2 9071 *0602 - *040 3 9070 *0803 - *0803 - *080 3 9066 *0803 - *080	-	9067	*0801	•	0404	۱ ۱	*0404	1040
3 9070 +0803 +030 3 9066 +0803 +060 9075 +0901 - +0101 - +030	N	9071	*0802	•	•	1	*0402	* 0404
.3 9066 *0803 - *0101 - *050	ب	9070	*0803	•	ı	1	*0301	*0501
9075 +0901 - +0101 - +030	س	9066	•0803	1	•	•	+0501	1000
		310	1060	1	*0101	•	.,,	* 0.0
CXXLC (0-0) +1001 + +1501	4	20.0					١	5

The allelic composition at DRB, DQA1 and DQB1 loci for the sequenced haplotypes corresponded to that expected according to published sequence information from well characterized homozygous cell lines unless indicated (*).

- Baplotypes carrying new allelic sequences (DRB1, DRB3, DQA1 or DQB1 loci).
- Only the tested heterozygote combinations are listed. The remainder of the 40 subjects tested were homozygotes or carried the haplotypes listed in this table.
- This DRB specificity (DR5x6) has been given this arbitrary designation according to serological, RFIP and sequence information.

Amplification and Direct Sequencing of DOA1 and DOB1 CDRAS in Subjects of Unknown HIA Type

DNA sequences have been determined for most HLA Class II allelic specificities defined by conventional 5 HLA typing techniques (March, S.G.E., Bodmer, J.G. HLA-DRB nucleotide sequences, 1990. Immunogenetics 31:141, 1990; Todd, J.A., Bell, J.I., McDefvitt, H.O.: HLA-DQB gene contributes to susceptibility and resistance to insulin-dependent diabetes mellitus. Mature 329:599, 1987). Comparisons of these sequences.

any given DQA1 or DQB1 homozygous or heterozygous allelic combination is characterized by a specific sequencing ladder.

Total RNA from PENNCs from 40 different

15 subjects was tested to evaluate if the allelic composition of DQA1 and DQB1 homo- and heterozygotes could be determined correctly by direct amplification and sequencing using Type 1 primers. These subjects had been previously secologically typed but the typing 20 information was not known to the investigator who

O information was not known to the investigator who assigned the Class II allelic specificities from the sequencing results. These 40 subjects comprised 27 different heterozygote combinations (Table III). All individuals were assigned DQA1 and DQB1 allelic

25 sequences that were consistent with the serological phenotypes. In all the heterozygotes tested, both alielic sequences could be read clearly from the composite sequence pattern. A unique pattern is found for every particular heterozygote combination in the

30 same way that certain RFIP banding patterns correspond to certain heterozygote allelic combinations. For instance, in a DQB2/DQB1.1 heterozygote one would find the sequence GGGG(A/T)T(T/A)CCGGGC(A/G) at codons 45 to 49 which can only be attributed to that particular

35 allele combination. In practice, interpretation of heterozygous sequence ladders is initiated by reading certain polymorphic positions where allele-specific bases may be found, such as, for instance, the second

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base of codon 46, where DQB1*0201 is the only allele that has an A. The sequences of the two possible templates are then deduced and compared with the sequences of all known alleles at the different loci.

5 In Figure 3 we show the overlapping ladder corresponding to a DQB1*0201/DQB1*0302 heterozygote; interpretation of the pattern is indicated on the side of the ladder.

The absence of expected bands or the presence

of unexpected bands for a particular allele or allelic combination is therefore suggestive of sequence heterogeneity, i.e., new alleles. The same can be said for DPA1 and DPB1 typing when appropriate primer combinations are used (Table III). For instance, substitution of the A at the second base of codon 46

15 would strongly suggest the presence of a sequence variant of DQB1+0201. Once detected, the sequence of the variant can be confirmed after selective amplification of the variant or by subcloning the amplified products.

 Amplification and Direct Sequencing of DRB cDWAS From Subjects of Unknown HLB Type

As described above, the use of Type 1 primers

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allows the unambiguous sequencing of all heterozygous combinations of DQA1 and DQB1 alleles. The same can be said for DPA1 and DPB1 typing when appropriate primer combinations are used (Table II). Because of the isotypic complexity of DRB genes (expression of more than one DRB locus by certain haplotypes), amplification and sequencing of cDNAs from DRB heterozygotes with Type 1 primers can generate up to four overlapping ladders, thus generating complex sequencing patterns.

above for DQA1 and DQB1 genes were amplified and 35 sequenced using DRB-specific Type 1 primers. As mentioned above, these 40 individuals comprised 27 different heterozygote combinations, including several examples from each of the groups of complex DRB allelic

DRB cDNAs from the same 40 individual tested

combinations which would generate up to four sequencing ladders. The DRB sequence ladders generated with Type 1 primers were analyzed as described above for DQA1 and DQB1 loci; highly polymorphic positions were analyzed first for the presence of bands unique to reconstitute.

- 5 first for the presence of bands unique to specific alleles or groups of alleles (i.e., DR4) and the sequences deduced and compared with the sequences of all known alleles at all loci. As example, in Figure 3 we show the ladder generated by sequencing a complex DRB
- 10 heterozygote (four overlapping ladders); the positions with two or more bands are indicated on the side of the Figure and assigned to each of the allelic types composing the complex sequencing pattern. For all but one sample, the information deduced from these
- 15 sequencing experiments matched the independently determined serological phenotypes of the subject under study as well as the DQA1 and DQB1 allelic types assigned to these individuals by direct sequencing of these games as described above. The inconsistent sample
- 20 had been serologically typed as DRw13/DR# but was typed by sequence analysis as DRw13/DRw8-Dw8.1. The presence of a DRB1*0801 allele instead of a DRB1*0401 allele was confirmed in a repeated experiment; we thus believe that the serological typing was in error. In all the 40
- 25 cases, all DQB1, DQA1 and DRB1 templates had been equally amplified and sequenced with a similar efficiency by the use of Type 1 primers. DRB3, DRB4 and DRB5 sequence ladders could be read in all but one case (a DRB3*0101 [DRw52a] sequence was not initially
- 30 observed in a DRw13/DRw17 heteroxygote). Since DRB3*0101 is in linkage disequilibrium with DRB1*0301, the former allele was expected to be found in the overlapping ladder as well. In order to rule out the possibility of an error, the investigator assigning the
- 35 HLA types from the sequencing ladders repeated the typing of this individual; the DRH3*0101 could be read in the repeated experiment.

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Although the results generated by the use of Type 1 primers were compatible with the serological phenotypes, the exclusive use of Type 1 primers will not allow in all cases to assign each of the specific

- 5 ladders to each of the expressed loci in all possible heterozygotes. Given below are the most complex situations which cannot be addressed by the exclusive use of Type 1 primers: 1) distinction among the different DR4 allelic sequences in certain heterozygotes 10 since they differ by only a few nucleotide base pairs and such differences could be masked by the presence of additional ladders; 2) to distinguish between DRB1*1601
- additional ladders; 2) to distinguish between DRB1*1601 and DRB1*1502 since their sequence differences will be masked by those of their linked DRB5 alleles; 3) to distinguish between DRB1*1301 and DRB1*1302 (which only differ at codon 86 since this difference can also be masked by other ladders; and finally 4) distinction between DRB1*0301 and DRB1*0302 in specific heterozygote combinations.
- 20 We have thus developed a more informative strategy to deal with DRB; this strategy, which consists of the additional use of non-conserved (Type 2) primers permits the clear elucidation of even the most complex combination of the four DRB sequences that might be
- 25 present in an individual. These non-conserved primers, as opposed to allele-specific primers, are designed to be used in reactions performed simultaneously with the reactions using Type 1 primers and aim at selectively amplifying certain ladders from the complex sequencing patterns without requiring previous typing information.
- analysis of the sequence variability of the second exon of the DRB genes has allowed us to identify two regions which could be used to design non-conserved (Type 2) primers: 1) codons 5-13; and 2) codons 29-35.
- 35 The sequence of the former region follows a groupspecific sequence pattern, i.e., a sequence shared by groups of alleles at individual loci. The later region

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10 IV). DRB templates, the type of templates amplified of mismatches between these primers and the different among the known alleles for each locus) (Tables I and moderate polymorphism (from 1 to 5 nucleotides different the latter two primers annealing to the second region of DRw11-, DRw12-, and DRw8- DRB1 ladders); 3) DRB25 DRB24 (specific for DRw17-, DRw18-, DRw13-, DRw14-, regions: 1) DRB23 (specific for DR2-DRB1 ladders); 2) DRB3, DRB4 and DRB5 genes. We designed five different (specific for DR4- DRB1 ladders); 4) DRB16 and 5) DRB17, non-conserved primers annealing to these two polymorphic exhibits a scattered nucleotide polymorphism in DRB1 and Because of the different nature and distribution

TABLE IV Contribution of Nucleotide Base Pair Mismatches Between 5' Amplification Primers and DRB Alleles to the Selective Amplification of Allelic And/or Non-Allelic DRB Transcripts

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specific combinations of primers used for the selective amplification of DRB transcripts.

the possible cDNAs that are required to obtain such

CDNA/PCR/sequencing reactions are shown in Table II

The results of this analysis are shown below

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with the sequences of the known DRB alleles at the

sequences of these primers carry from 0-12 mismatches results for DRB typing. Furthermore, since the which combination would give the best discriminatory We therefore tested these primers in order to determine

different DRB loci, their use allowed us to determine

the number of mismatches between the primers and each of

20

DRB4 and/or DRB5 loci in most heterozygote combinations.

amplification of certain transcripts from DRB1, DRB3, primers DRB16 and DRB17 will allow the random selective

DRB3, DRB4 or DRB5 cDNAs. On the contrary, the use of

cDNAs in any given heterozygote and will not amplify any

the first three primers will amplify up to two DRB1 selectively by these primers will be different. Each of

					DRB1				
	*0101-3	*1501	*1601-2 *1502	*1401- *0301/13		DR5×6## 8 *1101-4	*0801-3	*1201	*0701
RB16	0	2	1	4	3	2	3	5	4
ORB17	4	4	5	0	3	2	3	ī	3
RB23	5	0	. 0	7	4 '	4	7	8	8
RB24	6	5	5	0	4	D	0	0	12
RB25	6	4	4	8	, 0	В	8	8	5
		0R B1			t pre3/p	RB4/DRB5			
	*090	1 *10	001 D	RB5#	DRB3*0101	DRB3*0201	DRB3*03	101 pi	RB4
ORB16	1	:	3 · 4	3	4	5	4		2
DRB17	5		4	1	0	1	ō		2
RB23	2		4	6	5	5	5		3
DRB24	5		4	6	5	5	5		3
DRB25	5		4	В	6	5	5		4

DRB5 gene from cell line AMAI has an additional nucleotide substitution in the first of codon 30, in comparison with DRB5 genes of other DR2 haplotypes.

The DRB1 gene of this specificity (DR5x6) has been given this arbitrary designation according to serological, RFLP and sequence information.

TABLE V

Selective Amplification of DRB and DQB1 cDMA. In Combinations of Alleles Missatched with Type 2 Oligonucleotides(f)

DQ36	DQB1*0201	DQB1*0201/DQB1*0502
DQB6***	DQB1*0201/DQB1*0302	DCB1#0201/DCB1#030Z
DQB14	DQB1*0501	101041800/102041804
DQB14	DQB1*0501	DQB1*0603/DQB1*0101
DQB6	DQB1*0603	DQB1*0603/DQB1*0101
DQ86	DQB1+0301	DQB1+0301/DQB1+0502
DQB15	DQB1*0604	DQB1*0604/DQB1*0301
DQB6	DQB1*0201	DQB1*0201/DQB1*0603
DQB14	DQB1*0501	DQB1*0301/DQB1*0101
DQB6	DQB1*0301	DQB1*0301/DQB1*0101
ДОВ6	DQB1*0504	DQB1*0604/DQB1*0502
DREL6	DRB4+0101	DRB1+0401, DRB4*0101
DREIS	DRB1*1602	DRB1*1602,DRB5*0202
DRB17	DRB5*0201	DRB1*1601, DRB5*0201
DRB16	DRB1 *1601	DRB1*1601, DRB5*0201
DRB16**	DEB1*1601 + DEB1*0401	DRB1*1601, DRB5*0201/DRB1*0401, DRB4*0101
DRB17	DRB1*0401 + DRB5*0201	DRB1*1601, DRB5*0201/DRB1*0401, DRB4*0101
DRB16		DRB1*1501, DRB5*0101
DRB16	DREI+1101 + DR5x6	DR516, DRB3*0101/DRB1*1101, DRB3*0201
DRB16	+	DRB1*0405, DRB4*0101/DRB1*0301, DBB3*0101
DBB17	DRB1*1201 + DRB3*0201	DRB1*1201, DRB3*0201/DRB1*1101, DRB3*0201
DRB16	DRB1*1101	DRB1*1201, DRE3*0201/DRB1*1101, DRB3*0201
DRB17*	DRB5+0101	DRB1*1101,DRB3*0201/DRB1*1501,DRB5*0101
DRB16	DRB3*0201	DRB1*1101, DRB3*0201/DRB1*1501, DRB5*0101
DRB16	DRSIG	DR5x6, DRB3*0101/DRB1*0301, DRB3*0101
DRP17	DRES*0101	DR5x6, DRB3*0101/DRB1*08011
DEB16	DR81*0801	DR5x6, DRB3*0101/DRB1*0801
DRB16	DRB1+1601	DEB1+0301, DEB3+0101/DEB1+1601, DEB5+0201
DRB16	T080 + 1880	DRB1*1301,DRB3*0101/DRB1*0801
DRB16	DRB1*1601	DRB1*1301,DRB3*0101/DRB1*1601,DRB5*0201
DO Primer	Selected Alleles	Replotypes

³ combinations lacking those aliels the primers are fully matched with, for reasons of simplicity (see Table IV). Whenever these primers were used in hererozygotes carrying the alleles they specifically recognize, these alieles were selectively amplified. Note that in the examples shown in the Table the non-conserved In this Table we only show representative examples of haplotypic shown under "haplotypes". The selected alleles and the primers used primers selectively amplified the templates closer in sequence to the primer. The DRB and DQB1 alleles composing these haplotypes are are indicated in the two other columns. More than one individual

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Table. was tested for some of the heterozygote combinations listed in this

- DRB5*0101 and DRB3*0201 templates both have one mismatch with primer DRBII7. Selection of DRB5*0101 could be related to the differential positioning of the mismatch with respect to the primer.
- * A Veaker DRB4*0101 template was also observed
- Despite the presence of a mismatch between these two DQBI alleles. primer DODQB6 was not able to select either of them.

10 Ċ primers could differentially amplify DRB transcripts from the as few as one nucleotide substitution, provided that high combinations of allelic cDNAs that differ from each other in shown in Table V and Figure 4. As shown in Table V, Type 2 in heterozygotes not carrying the alleles specifically amplified in the PCR. Specific examples of the latter are fewest base pair mismatches with the primers were selectively recognized by the primers, the DRB templates which had the are matched with, these alleies were selectively amplified; tested. In heterozygotes carrying the alleles these primers certain DRB templates in all the heterozygous combinations These primers were able to selectively amplify

않 20 ļ, combination DRw13/DRw8-Dw8.1, the DRB1*0801 allele (3 possible that the differential positioning of the mismatches primer DRB17, this oligonucleotide selected the DRB5*0101 mismatches with the primer) was selected over DRB1*1301 and has an influence on the stability of the primer/cDNA complex within the sequence recognized by the oligonuclectide also sequence in a DRw11/DRw15 heterozygote (Table V). DRB3*0201 and DRB5*0101 genes all harbour one mismatch with the DRB16 oligonuclectide primer. Although DRB3*0101 or DRB3*0101 genes (each has 4 mismatches with the primer) by (annealing at 55°C). For example, in the heterozygote

stringency annealing conditions are used for the PCR

and hence on the outcome of the PCR.

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The ability of non-conserved primers to select certain alleles in heterozygote combinations was also tested for DQB1 genes (Table V). As with DRB-specific Type 2 primers, the use of high temperatures (55°C) in the annealing

- 5 step of the PCR was required for achieving the selective amplification of single DQB1 alleles in heterozygotes with non-conserved primers. For instance, when ennealing of primer DQB6 was allowed to proceed at 37°C in cDNAs from a DQB1*0301/DQB1*0501 and a DQB1*0201/DQB1*0603 heterozygote,
- 10 both alleles in both heterozygotes were equally amplified.

 At 55°C, the allele with the most homologous sequence to the
 5' primer, was amplified over the other in the PCR.

 Combinations of alleles both differing from the primer in two nucleotides but in different relative position were also
- 1.5 differentially amplified with a non-conserved primer. For instance, primer DQB6 selected the DQB1*0604 sequence in a DQB1*0604/DQB1*0502 heterozygote (Table V). Five nucleotides separate the two mismatches between the DQB1*0604 allele and the DQB6 primer, whereas only two nucleotides separate the
- 20 mismatches between the DQB1+0502 and the primer,

These results clearly indicate that the oligonucleotide primers annealing to polymorphic regions at the 5' end of the target cDNAs can be tailored to achieve a reproducible selective amplification of a limited number of

- 25 DRB or DQB templates in complex heterozygous combinations. Although the use of Type I primers allows the unambiguous sequencing of all possible DQA, DQB, DPA and DPB heterozygotes, such an approach will not give absolute discriminatory information for all DRB heterozygotes. We
- 30 have shown that the simultaneous use of Type 1 and Type 2 primers for DRB will permit the clear elucidation of even the most complex of all DRB heterozygote combinations. When DRB-SET is used for typing purposes, we perform three Type 2-reactions (using DRB23, 24 and 25) simultaneously with a Type
- 35 1-reaction (Table II). The simultaneous use of these

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reactions using these primers has the highest discriminatory power for complete DRB typing in a single run and allows the identification of novel sequence heterogeneity. Only one Type 1 reaction is required for DQB1, one for DQB1, one for DQB1 and one for DPB1 (Table II).

EXAMPLE III

<u>Petermining Unknown HLA Type of Subjects by Direct Sequencing of the Second Exon of Class II Genes</u>

individuals for sequence polymorphisms can be performed by the use of the methodology reported here which can also identify previously unknown allelic variants. Figures 2A and 2B show a flow-chart for the routine protocol used to 15 determine sequence allelism of individuals of unknown HLA types.

Employment of Primer Combinations for CDNA, PCR and Direct Sequencing Using RNA as Initial Template

invention provides single strand DNA anti-sense oligonucleotide primers that anneal to conserved regions of the gene mRNAs to be reverse transcribed, amplified and sequenced. These oligonucleotide primers include an

- region (codons 105 through 111) shared by all the alleles at all the DRB loci, the latter being DRB1, DRB3, DRB4 and DRB5, respectively (e.g., primer DRB20). Four simultaneous cDNA reactions (one per tube) are performed for DRB typing, all using primer DRB20 (reactions A, B, C and D in Table II and Figure 2A); (2) anneals to a conserved region (codons 105 through 111) shared by all the alleles at the DQB locus (e.g. primer DQB7) (reaction E in Table II and Figure 2A); (3)
- anneals to a conserved region (codons 147 through 157) shared 55 by all the alleles at the DQA locus (e.g. primer DQA9)

u oligonuclectides added to each of these reactions once the products are indicated below as well as in Table II and in cDNA synthesis is done in order to emplify and sequence the DPA19) (reaction I in Table II and Figure 2A). The specific 2A); (6) anneals to a conserved region (codons 222 through 228) shared by all the alleles at the DPA locus (e.g. primer locus (e.g. primer DPA14) (reaction H in Table II and Figure (codons 104 through 110) shared by all the alleles at the DPA alleles at the DPB locus (e.g. primer DPB11) (reaction G in Table II and Figure 2A); (5) anneals to a conserved region conserved region (codons 105 through 111) shared by all the (reaction F in Table II and Figure 2A); (4) anneals to a

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20 15 to DRB genes took place. expressed DRB loci of each chromosome (DRB1 and DRB3 or DRB4 four tubes where the cDNA synthesis reactions corresponding to -26 (e.g. oligonucleotide DRB11) is added to one of the conserved oligonuclectide primer which anneals to codons -32 or DRB5, depending on the haplotype -isotypic complexity-), a To amplify cDNA molecules corresponding to each The combination of the cDNA

25 different non-conserved oligonucleotides (also called Type 2) primer DRB24), 6-13 (e.g. primer DRB25), respectively. Each annealing to codons 7-13 (e.g. primer DRB23), 5-11 (e.g. expressed by a given individual. Each of the remaining three primer is used to amplify all the alleles at all DRB loci synthesis reaction primer and the newly added conserved tubes containing DRB cDNA products receives one of three

30 of cDNAs corresponding to different groups of alleles at the or two for each of the parental chromosomes). by these four reactions allows complete and accurate DRB1 locus. Comparison of the sequencing ladders generated four possible DRB genes expressed by a given individual (one interpretation of the sequences corresponding to each of the

non-conserved primer is designed to favor the amplification

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Çn parental chromosome). In the case of the DQA1 locus, a conserved single strand DNA oligonucleotide primer useful for genes expressed in any given individual (one for each primer DQB13) can be used for amplifying each of the DQB1 primer which anneals to codons 1-7 of the DQB cDNAs (e.g. For the DQB1 locus, a conserved oligonucleotide

ij oligonucleotide (e.g. primer DPB10) annealing to codons -19 primer DQA10). For the DPB1 locus, a conserved subject anneals to codons -10 to -4 of the DQA1 cDNA (e.g.

amplifying each of the DQA1 genes expressed in any given

oligonucleotide (e.g. primer DPA15) annealing to codons -23 to -17, is used to amplify each of the expressed DPA1 genes in any given subject. For DPA1 locus, a conserved to -13, is used to amplify each of the expressed DPB1 genes

20 15 gene. in a given subject. In a separate reaction, conserved primer reaction is targeted at a second polymorphic region of this expressed DPA1 genes in any individual. This second DPA1 DPALS, annealing to codons 59-65 of the DPAL cDNAs is used in combination with the cDNA primer DPA19 to amplify each of the

anti-sense oligonucleotide primer (e.g. DRB12) annealing to chain reaction products corresponding to DRB loci include an codons 87-94 of all the alleles at DRH loci; this primer is Primers useful in direct sequencing the polymerase

25 chain reaction products generated with the other three DRB used for sequencing the products generated by the first of 97-103 of all the alleles at DRBI locus can be used (e.g. reactions, an anti-sense oligonucleotide annealing to codons the four DRB reactions. For direct sequencing the polymerase

80 of downstream polymorphic regions of DRB1 genes not seen in oligonucleotide in these three DRB reactions allows reading primer DRH30). The use of a different sequencing primer DRB12. the first DRB reaction which uses the example sequencing Primers useful in direct sequencing the

ដូ polymerase chain reaction products corresponding to DQBI

locus include an anti-sense oligonucleotide primer (e.g. DQB5) annealing to codons 78-83 of all the alleles at this locus. Direct sequencing of polymerase chain reaction products corresponding to DQA1 locus include an anti-sense oligonucleotide primer (e.g. DDA9) annealing to codens on oligonucleotide primer (e.g. DDA9) annealing to codens out

- 5 oligonucleotide primer (e.g. DQA29) annealing to codons 88-95 of all the alleles at this locus. Direct sequencing of polymerase chain reaction products corresponding to DPB1 locus include a sense oligonucleotide primer (e.g. DPB13) annealing to codons 12/-5 of all the alleles at this locus.

 10 For direct sequencing of polymerase chain reaction products for the DPA1 reaction which used primers DPA14 and DPA15, an anti-sense oligonucleotide annealing to codons 88-94 of all the alleles at this locus can be used (e.g. primer DPA16).
- For direct sequencing of polymerase chain reaction products for the DPA1 reaction which used primers DPA19 and DPA18, an anti-sense oligonuclectide annealing to codons 214-220 of all the alleles at this locus can be used (e.g. primer DPA20).

Employment of Primer Combinations for PCR and Direct Sequencing Using DNA Templates

To amplify DNA molecules corresponding to each DRB loci of each chromosome a conserved anti-sense oligonucleotide primer annealing to base pairs 18-38 of intron 3 (e.g. oligonucleotide DRB1406) is added to each of four PCR reaction tubes (reactions S, V, T and U in Table II and Pigure 2B). Each of these four tubes will receive a different additional oligonucleotide annealing to codons -4 to +3 (e.g. primer DRB22), to codons 7-13 (e.g. primer

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DRB23), 5-11 (e.g. primer DRB24), 6-13 (e.g. primer DRB25),

30 respectively. The first reaction is used to amplify all the
alleles at all DRB locd carried by a given individual. Bach
of the remaining three reactions is designed to favor the
amplification of DNA corresponding to different groups of
alleles at the DRB1 locus. As with RNA templates, comparison

35 of the sequencing ladders generated by these four reactions

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Primers useful in direct sequencing the polymerase chain

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allows complete and accurate interpretation of the sequences corresponding to each of the four possible DRB genes expressed by a given individual (one or two for each of the parental chromosomes).

20 15 ö W in Table II and Figure 2B). For the DPB1 locus, two Figure 2B). genes carried by a given subject (reaction I in Table II and pairs 55-71 of intron 3) are used to amplify each of the DPA1 pairs -69 to -50 of intron 2) and DPA11 (annealing to base conserved oligonucleotide such as DPA10 (annealing to base (reaction X in Table II and Figure 2B). For DPA1 locus a amplify each of the DPB1 genes carried by any given subject DPB15, annealing base pairs 39-59 of intron 3) are used to to base pairs -42 to -62 of intron 2, and a primer e.g. conserved oligonuclectides (a primer, e.g. DPB14, annealing given individual (one for each parental chromosome) (reaction used for amplifying each of the DQB1 genes carried by any 11-17 (e.g. primer DQB931) or 1-7 (e.g. primer DQB13) can be primers which anneal to codons 88-94 (e.g. primer DQB932) and For the DQB1 locus, two conserved oligonucleotide

뚕 25 DRB1400). The use of a different sequencing oligonucleotide of all the alleles at DRB1 locus can be used (e.g. primer reaction products generated with the other three DRB of all alleles at DRB loci; this primer is used for oligonucleotide primer (e.g. DRB12) annealing to codons 87-94 corresponding to DRB loci include an anti-sense polymorphic regions of DRB1 genes not seen in the first DRB in these three DRB reactions allows reading of downstream reactions, a sense oligonucleotide annealing to codons 39-46 DRB reactions. For direct sequencing the polymerase chain chain reaction products generated from DNA templates reaction which uses the example sequencing primer DRB12. sequencing the products generated by the first of the four Primers useful in direct sequencing the polymerase

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reaction products corresponding to DQBI locus include an anti-sense oligonuclectide primer (e.g. DQBS) annealing to codons 78-83 of all the alleles at this locus. Direct sequencing of polymerase chain reaction products corresponding to DPBI locus include an anti-sense oligonuclectide primer (e.g. DPBI6) annealing to base pairs 1-21 of intron 3 of all the alleles at this locus. For direct sequencing of polymerase chain reaction products for the DPAI reaction an anti-sense oligonuclectide annealing to codons 76-82 of all the alleles at this locus can be used (e.g. primer DPAI2).

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Procedure for Determining Unknown HLA Type

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A subject of unknown HLA type, diseased or not, is to be typed for Class II HLA polymorphism. From 10 to 50 mL of peripheral blood are drawn. The peripheral blood mononuclear cells are prepared by centrifugation over Ficoll-Hypaque gradients. The cells are then lysed in guanidium isothyccianate and total cellular RNA prepared using

20 conventional methods (either by centrifugation on cesium chloride gradients, which lasts about 16 hours, or by the guanidium isothyocianate-phenoi-chlorophorm extraction method, which can be performed in less than 4 hours. See Gounh, supra (1988); Johns et al., Anal, Biochem., 180:276

25 (1989). Otherwise genomic DNA from these cells or other sources (hair, blood stains, sperm, etc.) can be prepared with conventional methods such as provided by Higuchi, R. in PCR Technology, Erlich, M. (ed.), Stockton Press:31 (1989). DQB1, DQA1, DRB (DRB1, DRB3/4/5), DPA1 and DPB1 cDNA

30 molecules are synthesized from total RNA using locus-specific primers. Approximately, one microgram of RNA is reverse transcribed with MolVRT (reverse transcriptase) and DRB (CODRB20), DQB (CODQB7), DQA (CODQA9), DPB (DPB11) and DPA (DPA14, DPA19) (optional) -specific non-sense primers in a 20 35 ut final volume reaction (30-60 minute incubation). The

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reaction for each Class II gene is performed in a different tube, but they can be performed in the same tube if preferred. For routine purposes, four simultaneous reactions are performed for DRB, one for DQB, one for DQA1, one for DPB1, and two for DPA1 gene products.

Once these reactions are completed, the enzymatic amplification of the respective cDNA molecules is then performed by directly adding to the 20 uL reverse transcription reaction, the reagents needed for the 10 amplification step. Alternatively, if DNA is used, the primer combinations used for the PCR are those shown in Table II herein (the anti-sense primers as well as the sense primers will be different). This includes the PCR reagents

15 primers. This example uses four reactions for DRB (tubes 1, 2, 3 and 4), one for DQB (tube 5), one for DQA (tube 6), one for DPB (tube 7), and two for DPA (tubes 8 and 9, respectively). Reactions 2, 3 and 4 incorporate primers DRB23, DRB24 and DRB25, respectively. For rapid typing (in combinations. Alternative combinations of the primers that can be used are shown in Table II.

and appropriate conserved and non-conserved oligonuclectide

about 15 minutes using Centricon (Amicon, Ultrafree (milli25 pare)) or similar columns to remove unincorporated primers
and dNTPs. The retentate or one half of the recovered
retentate for each reaction is then directly sequenced using
Tag polymerase and the primers described in Table II for each
combination of primers used in the cDNA/PCR reactions using
9-32 end-labeled (10 minutes) locus-specific sequencing
primers (35 minutes).

Once completed, the reactions are spun-dialyzed for

The sequencing reactions products are loaded on an acrylmide gel, electrophoresed in 2-3 hours and exposed to X-ray films for 4-12 hours. The gels are read and results from

all possible alleles. gels are compared to nuclectide sequences corresponding to

the nucleotide sequences of all alleles of all haplotypes and allelic sequences. routines which indicate how the comparison is to be performed as well as subroutines which will allow identification of new or using a personal computer and a software package including Comparisons can be made visually using the maked eye

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

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	(X;		(viii)		(vi)		(ব		(¥¥)	(1111)	(ii)		(±)
(A) TELEPHONE: (612) 332-5300 (B) TELEPAX: (612) 332-9081	TELECOMMUNICATION INFORMATION:	(A) NAME: KOWAIChyk, Alan W. (B) REGISTRATION NUMBER: 31,535 (C) REFERENCE/DOCKET NUMBER: 600.243WO	ATTORNEY INFORMATION:	(A) APPLICATION NUMBER: 07/665,960 (B) FILING DATE: 06-MAR-1991 (C) CLASSIFICATION:	CURRENT APPLICATION DATA:	720 Kb. COMPUTER: Northgate 386 ODERATING SYSTEM: DOS 4.9 SOFTWARE: WordPerfect 5	COMPUTER READABLE FORM: (A) MEDIUM TYPE: Diskette. 3.5 inch.	(A) ADDRESSEE: Merchant & Gould (B) STREFT: 3100 Norwest Center (C) CITY: Minneapolis (D) STAFE: Minnesota (E) COUNTRY: USA (F) ZIP: 55402	CORRESPONDENCE ADDRESS:	NUMBER OF SEQUENCES: 49	TITLE OF INVENTION: DNA Sequence-Based HLA Typing Method		APPLICANT: Santamaria, Pedro

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(2) INFORMATION FOR SEQUENCE ID NO: 2: (2) INFORMATION FOR SEQUENCE ID NO: 1: ORIGINAL SOURCE: 3 FEATURE: G GTG GTP GAG GGC CTC TGT CC FRAGMENT TYPE: Internal Fragment ANTI-SENSE: yes MOLECULE TYPE: Genomic DNA **3**60€ SEQUENCE CHARACTERISTICS: SEQUENCE DESCRIPTION: SEQ ID NO: 1: (A) NAME/KEY: Oligonucleotide Primer DQB7 **B** FRAGMENT TYPE: Internal Fragment MOLECULE TYPE: Genomic DNA ORIGINAL SOURCE: ANTI-SENSE: yes SEQUENCE CHARACTERISTICS: NAME/KEY: Oligonucleotide Primer DR920 LENGTH: 21 base pairs TYPE: Nucleic Acid STRANDEDNESS: Single TOPOLOGY: Linear LOCATION: Anneals to codons 105 to 111 of the DQB1 transcript of MLA class II LENGTH: 21 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single
TOPOLOGY: Linear Synthetically Derived Synthetically Derived PCT/US92/01675 21 WO 92/15711 S ŝ 40 35 8 25 20 15 10 (2) INFORMATION FOR SEQUENCE ID NO: 4: (2) IMPORMATION FOR SEQUENCE ID NO: 3: Ξ (xì) (EX (Ŧž) (4¥) (±±) **₫** E (X: 808 808 SEQUENCE CHARACTERISTICS: GGT GAG GTT ACT GAT CTT GAA G SEQUENCE DESCRIPTION: SEQ ID NO: 3: (B) LOCATION: Anneals to codons 148 to 155 of the DQA1 transcript of HLA class II FEATURE: ORIGINAL SOURCE: (A) NAME/KEY: Oligonuclectide Primer DQA9 FRAGMENT TYPE: Internal Fragment ANTI-SENSE: yes MOLECULE TYPE: Genomic DNA **BOB** SEQUENCE CHARACTERISTICS: ere ere exe eee ere eer err SEQUENCE DESCRIPTION: SEQ ID NO: 2: (B) LENGTH: 21 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single
TOPOLOGY: Linear LENGTH: 22 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single
TOPOLOGY: Linear LOCATION: Anneals to codons 105 to 111 of the DRB1, DRB3, DRB4 and DRB5 transcripts of HLA class II -54-Synthetically Derived PCT/US92/01675

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(B) LOCATION: Anneals to codons -4 to +3 of the DRB1, DRB3, DRB4 and DRB5 transcripts of HLA class II	(ix) FEATURE:(A) NAME/KEY: Oligonucleotide Primer.DBR22	ANTI-SENSE: no FRAGMENT TYPE: Internal ORIGINAL SOURCE: Syr	(A) IENGTH: 21 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear (ii) MOLECULE TYPE: General C DNA	(2) INFORMATION FOR SEQUENCE ID NO: 5: (1) SEQUENCE CHARACTERISTICS:	(x1) SEQUENCE DESCRIPTION: SEQ ID NO: 4: AGA GAC TOT CCC GAG GAT TTC 21 Arg Asp Ser Pro Glu Asp Phe 1 5	(A) NAME/KEY: Oligonuclectide Primer DQB13 (B) LOCATION: Anneals to codons 1 to 7 of the DQB1 transcript of HLA class II	(1) ANTI-SENSE: no (v) FRAGMENT TYPE: Internal Pragment (vi) ORIGINAL SOURCE: Synthetically Derived (ix) FRATURE:	
(A) (AT) PAREL (AT) (B) (C) (B) (C) (B) (C) (C) (C) (C) (C) (C) (C) (C) (C) (C	45 (2) INFORMATION FO	(xi) sect	30 (B)	25 (vi) ORIG (ix) FEAT	$\langle \widetilde{\mathbf{p}} \rangle$ $\langle \mathbf{p} \rangle$	10 (2) INFORMATION F (1) SEQ (A) (B)	5 CTG	(xi) SEQ
(A) LENGTH: 20 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear MOLECULE TYPE: Genomic DNA ANTI-SENSE: no	N FOR SEQUENCE ID NO: 7: SEQUENCE CHARACTERISTICS:	SEQUENCE DESCRIPTION: SEQ ID NO: 6: TG TTC TCC AGC ATG GTG TGT C 21 Phe Ser Ser Met Val Cys Leu -30	DRB11 LOCATION: Anneals to codons -33 to -26 of the DRB1, DRB3, DRB4 and DRB5 transcripts of HLA class II	ORIGINAL SOURCE: Synthetically Derived FRATURE: (A) NAME/KEY: Oligonucleotide Frimer		N FOR SEQUENCE ID NO: 6: SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single	CTG GCT TTG GCT GGG GAC ACC Leu âla Leu âla Gly Asp Thr -1 1	-56- SEQUENCE DESCRIPTION: SEQ ID NO: 5:

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TOG COT OTG CAG GOT CGC GCG 21	SEQUENCE DESCRIPTION: SEQ ID NO: 8:	OI THE DOBS TRANSCRAPT OF HEA Class	(B) LOCATION: Anneals to codons 88 to 94	(A) NAME/KEY: Cligonucleotide Primer DQB932	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRAGHENT TYPE: Internal Fragment	ANTI-SENSE: yes	HOLECULE TYPE: Genomic DNA	(C) STRANDEDNESS: Single (D) TOPOLOGY: Linear			INFORMATION FOR SECTIONCE ID NO. 8.		:10	The The Val Met Se	SEQUENCE DESCRIPTION: SEQ ID NO: 7:		(H) LOCATION: Anneals to codons -10 to -4 of the DQA1 transcript of HLA class		FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRACMENT TYPE: Internal Fragment
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(v1)	(४)	(vì)	(11)			· (1)	(2) INFORMA				(xt)				(1x)	(¥i)	(খ	(iv)	(11)		Ŧ	()	(2) INFORMS
ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Eragment	ANTI-SENSE: yes	MOLECULE TYPE: Genomic DNA	STRANDEDNE TOPOLOGY:	(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid	SEQUENCE CHARACTERISTICS:	INFORMATION FOR SEQUENCE ID NO: 10:		15	TIT AAG GGC ANG TGC TAC TIC 21	SEQUENCE DESCRIPTION: SEQ ID NO: 9:	(B) LOCATION: Anneals to codons il to I7 of the DQB1 transcript of HLA class II		(A) NAME/KEY: Oligonucleotide Primer	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Fragment	ANTI-SENSE: no	MOLECULE TYPE: Genomic DNA	(A) LENGTH: 21 Dase pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear		SECURNCE CHARACTERISTICS:	(2) INFORMATION FOR SEQUENCE ID NO: 9:

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FRAGMENT TYPE: Internal Fragment

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(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: Nucleic Acid	ORMATIO		AGG ATA CAC AGT CAC CTT AGG 21	45 (x1) SEQUENCE DESCRIPTION: SEQ ID NO: 11:	of the DRB1, DRB3, DRB4 and DRB5 transcripts of HLA class II	40 (B) LOCATION: Anneals to codons 97 to 103	(A) NAME/KEY: Oligonuclectide Primer DRE30	35 (ix) FEATURE:	(vi) ORIGINAL SOURCE: Synthetically Derived	30 (v) FRACMENT TYPE: Internal Fragment	(iv) ANTI-SENSR: yes	(11) MOLECULE TYPE: Genomic DNA	(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDHESS: Single (D) TOPOLOGY: Linear	UENCE CHARAC	(2) INFORMATION FOR SEQUENCE ID NO: 11:	15	A TGG GGA GAT GGT CAC TGT GG 21	10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:	of the DQB1 transcript of HLA class	5 (R) LOCATION: Anneals to codons 97 to 104	(A)	(ix) PEATURE:	-591	WO 92/15711 PCT/US92/01675	
(B) LOCATION: Anneals to codons 8/ to 94 of the DRB1, DRB3, DRB4 and DRB5 transcripts of HLA class II 55	NAME/KEY: Oligonucleotide Primer DRB12	(ix) PRATURE:	45 (vi) ORIGINAL SOURCE: Synthetically Derived	(v) FRACMENT TYPE: Internal Fragment	_	(ii) MOLECULE TYPE: Genomic DNA	(C) STRANDEDNESS: Single (D) TOPOLOGY: Linear	(A)	(i) SEQUENCE CHARACTERISTICS:	30 (2) INFORMATION FOR SEQUENCE ID NO: 13:		25 GTA GTT GTG TCT GCA CAC 18	20 of the DQB1 transcript of HLA class II (x1) SEQUENCE DESCRIPTION: SEQ ID NO: 12:	(B) LOCATION: Anneals to codons 78 to 83	15	(A) NAME/KEY: Oligonuclectide Primer DQB5	(ix) FEATURE:	(V) FRAGMENT TYPE: Internal respnent Original Source: Synthetically Derived	(iv)	(ii) MOLECULE TYPE: Genomic DNA	(C) STRANDEDNESS: Single (D) TOPOLOGY: Linear	-60:		WO 92/15711 PCT/US92/01675	

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(x 1)		WO 92/15711
SECTION DESCRIPTION, SEC ID NO. 13.	-61-	
3 1		PCT/US92/01675

20 5 10 Ų (2) INFORMATION FOR SEQUENCE ID NO: 14: ₹ (±±) (i▼) (£ (XX) G. FRAGMENT TYPE: Internal Fragment MOLECULE TYPE: Genomic DNA ANTI-SENSE: Yes වීම **E**€ SEQUENCE CHARACTERISTICS: SECUENCE DESCRIPTION: SEC ID NO: CCG CTG CAC TGT GAA GCT C TYPE: STRANDEDNESS: Single TOPOLOGY: Linear 23 base pairs Nucleic Acid 1.0 20

25 (¥ž) Ê PEATURE: ORIGINAL SOURCE: Synthetically Derived

 $\widehat{\Xi}$ LOCATION: Anneals to codons 82 to 89 of the DQA1 transcript of HLA class

SEQUENCE DESCRIPTION: SEQ ID NO: 14:

(xi)

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CAC GGT TCC GGT AGC AGC GGT AG

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ORIGINAL SOURCE:

Synthetically Derived

FRAGMENT TYPE: Internal Fragment

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FEATURE:

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NAME/KEY: Oligonuclectide Primer DRB1400

(B)

LOCATION: Anneals to codons 38 to 45 of the DRB1, DRB3, DRB4 and DRB5 transcripts of HLA class II

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MOLECULE TYPE: Genomic DNA

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ANTI-SENSE: no

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SEQUENCE CHARACTERISTICS:

LENGTH: 20 base pairs TYPE: Nucleic Acid STRANDEDNESS: Single TOPDLOGY: Linear

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(2) INFORMATION FOR SEQUENCE ID NO: 16:

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SEQUENCE DESCRIPTION: SEQ ID NO: 16: 3 OGC TIC GAC AGC GAC GIG G Val Arg Phe Asp Ser Asp Val Gly 40

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FRAGMENT TYPE: Internal Fragment

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(ii)

MOLECULE TYPE: Genomic DNA

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(iv) ANTI-SENSE: no

(2) INFORMATION FOR SEQUENCE ID NO: 15: **B**0B5 SEQUENCE CHARACTERISTICS: (A) WAME/KEY: Oligonuclectide Primer DQA29 LENGTH: 18 base pairs TYPE: Nucleic Acid STRANDEDNESS: Single TOPOLOGY: Linear LENGTH:

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(¥. ORIGINAL SOURCE:

(x1) FEATURE:

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Ě SEQUENCE DESCRIPTION: SEQ ID NO: 15:

TAC GGT CCC TCT GGC CAG Tyr Gly Pro Ser Glu Gln 20

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LOCATION: Anneals to codons 19 to 24 of the DQA1 transcript of HLA class

 $\widehat{\mathbb{R}}$ NAME/KEY: Oligonucleotids Primer DQA30

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		(XT)				(ix)	(¥i)	₹)	(4T)	(ii)			(Ŧ)	NFORM				(JX)				(xì	(YY)	₹	(tv)	
	AAC CCC GTA GTT GTG TCT GCA 21	SEQUENCE DESCRIPTION: SEQ ID NO: 21:	rtanecifes of APV CISS II	(B) LOCATION: Anneals to codons 79 to 85 of the DRB1, DRB3, DRB4 and DRB5		FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Fragment	ANTI-SENSE: yes	Ŕ	(C) STRANDEDNESS: Single (C) TOPOTOGY: Finest		SEQUENCE CHARACTERISTICS:	INFORMATION FOR SEQUENCE ID NO: 21:			GCCAAGAGTG GGCCTCGCAG C 21	SEQUENCE DESCRIPTION: SEQ ID NO: 20;	DRB5 transcripts of MLA class II	(B) LOCATION: Anneals to bp18-38 to	(A) NAME/KEY: Oligonuclectide Primer DQB1406	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Fragment	ANTI-SENSE: no	-65-
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(x1)	(¥i)	(₹)	(iv)	(ii)		Ξ)		(2) INFORMAT			(xi.)					(x1)	(vi)	(¥)	(iv)	(H)			(±)	(2) INFORMA		
FEATURE;	ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Fragment	ANTI-SENSE: no	MOLECULE TYPE: Genomic DNA	(A) LENGTH: 21 DASE PAIRS (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear	UKNCK CHARAC		INFORMATION FOR SECURENCE ID NO: 23:	,	GGG GAC ACC CGA CCA CGT TTC 21 Gly Ala Thr Arg Fro Arg Phe 1	SEQUENCE DESCRIPTION: SEQ ID NO: 22:	transcripts of HLA class II	(B) LOCATION: Anneals to codons 1 to 7 of		(A) NAME/KEY: Oligonucleotide Primer DRB824	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Fragment	ANTI-SENSE: no	MOLECULE TYPE: Genomic DNA	CA STONE	(A) LENGTH: 21 base pairs	SEQUENCE CHARACTERISTICS:	INFORMATION FOR SEQUENCE ID NO: 22:		106.1

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(2) INFORMATION FOR SEQUENCE ID NO: 25: (2) INFORMATION FOR SEQUENCE ID NO: 24: $\widehat{\mathbb{E}}$ (XX) (XX) (Y1) 3 (A.) Ξ (i) (xi SEQUENCE CHARACTERISTICS: SEQUENCE DESCRIPTION: SEQ ID NO: 24: (A)97EENGTH: (B) TYPE: GET GTG GTG CAA GGG CCC (B) (<u>A</u>) FEATURE: ORIGINAL SOURCE: FRAGMENT TYPE: Internal Fragment ANTI-SENSE: yes SEQUENCE CHARACTERISTICS: CGG ACA GTG GCT CTG ACG GCG Arg Thr Val Ala Leu Tyr Ala -15 SEQUENCE DESCRIPTION: SEQ ID NO: 23: (A) MOLECULE TYPE: Genemic DNA (₿ IOCATION: Anneals to codons 105 to 111 of the DPB1 transcript of HLA class II NAME/KEY: Oligonuclectide Primer DPB11 NAME/KEY: Oligonucleotide Primer DPB10 TOPOLOGY: Linear IENGTH: 21 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single LOCATION: Anneals to codons -19 to -13 of the DPB1 transcript of HLA class II -67-21 base pairs Nucleic Acid Synthetically Derived 21 ភ 50 ŝ 6 35 30 25 20 5 5 (2) INFORMATION FOR SEQUENCE ID NO: 26: (4) (iv) (ii) <u>E</u> 3 $\widehat{\mathbb{E}}$ (¥i) (¥i) ₹ £ (i∀) (ii) (B) LOCATION: Anneals to codons -5 to +2 of the DPB1 transcript of HLA class II (A) NAME/KEY: Oligonuclectide Primer DPB13 FEATURE: ORIGINAL SOURCE: €ଞ୍ଚିତ୍ର FRAGMENT TYPE: Internal Fragment SEQUENCE DESCRIPTION: SEQ ID NO: 25: ANTI-SENSE: no MOLECULE TYPE: Genomic DNA SEQUENCE CHARACTERISTICS: (B) LOCATION: Anneals to codons 97 to 103 of the DPB1 transcript of HLA class II CITY GGA GGG GGA AAC AITY CAC (A) NAME/KEY: Oligonucleotide Primer DPB12 FEATURE: ORIGINAL SOURCE: FRAGMENT TYPE: Internal Fragment ANTI-SENSE: yes MOLECULE TYPE: Genomic DNA 90 LENGTH: 21 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single
TOPOLOGY: Linear STRANDEDNESS: Single TOPOLOGY: Linear

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(vi) ORIGINAL SOURCE: Synthetically Derived	(v) FRAGMENT TYPE: Internal Fragment	(iv) ANTI-SENSE: no	(ii) MCLECULE TYPE: Genomic DNA	TOPOLOGY:	(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid	(1) SEQUENCE CHARACTERISTICS:	(2) INFORMATION FOR SEQUENCE ID NO: 28:	AGAGGUBGAA AGAGGATTAG A	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:	HLA class II	(B) LOCATION: Anneals to bp-42/-46 to	Drate	(A) NAME/KEY: Oligonucleotide Primer	(ix) FEATURE;	(vi) ORIGINAL SOURCE: Synthetically Derived	(v) FRAGMENT TYPE: Internal Fragment	0 (iv) ANTI-SENSE: no	(ii) NOLECULE TYPE: Genomic DNA	5 (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear		(1) SEQUENCE CHARACTERISTICS:	(2) INFORMATION FOR SEQUENCE ID NO: 27:	5 -12 *5	CTG ATG GTG CTG CTC ACA	(x1) SEQUENCE DESCRIPTION: SEQ ID NO: 26	-69-1
rived 55		(2)	50		45		40	21	35	COX		J.	9r	25	erived	b	20	(2)	15		10		Un			
		INFORMATION FOR SECUENCE ID NO: 30:		CGGCCCAAAG CCCTCACTCA C 21	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:	3 of the DPB1 transcript of HLA class	(B) LOCATION: Anneals to bp1-21 to intron	(A) NAME/KEY: Oligonucleotide Primer DPB16	(1x) FEATURE:	(vi) ORIGINAL SOURCE: Synthetically Derived	(v) FRAGMENT TYPE: Internal Fragment	(17) ANTI-SENSE: no	(11) MOLECULE TYPE: Genomic DNA	(D) TOPOLOGY: Linear	(B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single	(A) LENGTH.	(i) SECTION THARAC) INFORMATION FOR SEQUENCE ID NO: 29:		S SOSOOOS ASSOSIONS	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:	(B) LOCATION: Anneals to bp39-59 to intron 3 of the DPB1 transcript of HLA class II	(A) NAME/KEY: Oligonucleotide Primer DPB15	(ix) FEATURE:		-70

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20 15 5 (2) INFORMATION FOR SEQUENCE ID NO: 31: (vi) (iv) (ii) (<u>x</u>) 3 (XX) Ë (ix) (AT) 3 (iv) (33) e Ð FEATURE: ORIGINAL SOURCE: FRAGMENT TYPE: Internal Fragment ANTI-SENSE: yes MOLECULE TYPE: Genomic DNA SEQUENCE CHARACTERISTICS: Ē 380 SEQUENCE DESCRIPTION: SEQ ID NO: 30: CECTCATETC CECCCCCTCC C **B** (A) NAME/KEY: Oligonucleotide Primer DPB17 PRATURE: ORIGINAL SOURCE: FRACMENT TYPE: Internal Fragment ANTI-SENSE: no MOLECULE TYPE: Genomic DNA ⊕<u>@</u> NAME/KEY: Oligonuclectide Primer DPA14 LOCATION: Anneals to codons 104 to 110 of the DPA1 transcript of HLA class II TOPOLOGY: Linear IOCATION: Anneals to bp-6/-26 to intron 2 of the DPB1 transcript of HLA class II LENGTH: 21 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single TYPE: Nucleic Acid STRANDEDNESS: Single TOPOLOGY: Linear -71-Synthetically Derived Synthetically Derived 21 S 50 ŝ 30 40 ü 25 20 15 ö (± Ξ (xi) (iv)

WO 92/15711 Œ SEQUENCE DESCRIPTION: SEQ ID NO: GTC AAT GTG GCA GAT GAG GGT 31: 21

(2) INFORMATION FOR SEQUENCE ID NO: 32:

(i) SEQUENCE CHARACTERISTICS:

LENGTH: 21 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single
TOPCLOGY: Linear

(iii) MOLECULE TYPE: Genomic DNA

ANTI-SENSE: yes

₹ FRAGMENT TYPE: Internal Fragment

(vi) ORIGINAL SOURCE: Synthetically Derived

Ė PEATURE:

(A)NAME/KEY: Oligonucleotide Primer DPA15

B LOCATION: Anneals to codons -17 to -23 of the DPA1 transcript of HLA class II

CAT ATC AGA GCT GTG ATC TTG

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SEQUENCE DESCRIPTION: SEQ ID NO: 32:

(2) INFORMATION FOR SEQUENCE ID NO: 33:

SEQUENCE CHARACTERISTICS:

LENGTH: 21 base pairs TYPE: Nucleic Acid STRANDEDNESS: Single TOPOLOGY: Linear

MOLECULE TYPE: Genomic DNA

(F ANTI-SENSE: yes

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₹ FRAGMENT TYPE: Internal Fragment

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ORIGINAL SOURCE:

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(2) INFORMATION FOR SEQUENCE ID NO: 35:

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			(1				(xi x)	(1 A	₹	(vì)	(11)		(±)	MFORMAT:			(XX)				(x ₇)
	Ala	CING CING AGT CINC CGA GGA GCT 21	SEQUENCE DESCRIPTION: SEQ ID NO: 34:	of the DPA1 transcript of HLA class	(B) LOCATION: Anneals to codons -3 to -9	(A) NAME/KEY: Oligonucleotide Primer DPA17	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Fragment	ANTI-SENSE: no	MOLECULE TYPE: Genomic DNA	(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear	Ę	INFORMATION FOR SEQUENCE ID NO: 34:		CIT GGG AAA CAC GGT CAC CIC . 21	D NO: 3:		(B) LOCATION: Anneals to codons 88 to 94 of the DPA1 transcript of HLA class	(A) NAME/KEY: Oligonucleotide Primer DPA16	FRATURE:
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	(ix)	(vi)	(∀)	(iv)	(11)			(±)	(2) INFORMATION		(xi			(ž	(vi)	(₹	(iv)	(II)			(1)
(A) NAME/KEY: Oligonuciectide Primer DPA11	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRACKENT TYPE: Internal Fragment	ANTI-SENSE: no	MOLECULE TYPE: Genemic DNA	(A) LENGTH 21 DASS PALES (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear		SEQUENCE CHARACTERISTICS:	ON FOR SEQUENCE ID NO: 36:	CTCTAGCTT GACCACTIGC 20	SEQUENCE DESCRIPTION: SEQ ID NO: 35:	(B) LOCATION: Anneals to bp-69/-50 of intron 2 of the DPA1 transcript of HLA class II	(A) NAME/KEY: Oligonuclectide Primer DPA10	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Fragment	ANTI-SENSE: no	MOLECULE TYPE: Genomic DNA	DEDNI		SEQUENCE CHARACTERISTICS:

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•		(B) LOCATION: Anneals to bp55-71 to intron 3 of the DDA1 transcript of			1 / 6 1
٠		HIA class II		(♥)	FRAGMENT TYPE: Internal Fragment
ហ	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 36:	л	(vi)	ORIGINAL SOURCE: Synthetically Derived
		AGTCTGAGGG TGGCAGAGAG G 21	Ĺ	(ix)	FEATURE:
10					(A) NAME/KEY: Oligonuclectide Primer DPA18
ລ) INFORMAT	(2) INFORMATION FOR SEQUENCE ID NO: 37:	10		
	(1)	SEQUENCE CHARACTERISTICS:			of the DPA1 transcript of HLA class
15		(A) LENGTH: 21 base pairs (B) TYPR: Nucleic Acid	5 .	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 38:
					CTG GCT AAC ATT GCT ATA TTG 21
20	(11)	MOLECULE TYPE: Genomic DNA	š		60 65
	(iv)	ANTI-SENSE: yes	ě		
J.	(v)	FRAGHENT TYPE: Internal Fragment			
ţ	(v1)	ORIGINAL SOURCE: Synthetically Derived	25 (z) enformati	(2) INFORMATION FOR SEQUENCE ID NO: 39:
	i e	י מתוחסי.		Ξ	SEQUENCE CHARACTERISTICS:
30	٤	(A) NAME/XEY: Oligonucleotide Primer DPA12	30		(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single
35		(B) LOCATION: Anneals to codons 76 to 82 of the DPA1 transcript of HLA class II	យ	(±i)	EN C
	<u> </u>		;	(iv)	ANTI-SENSE: yes
40	į	0 NO: 31		(4)	FRAGMENT TYPE: Internal fragment
:		ביח שמד בות חדו המני טרים 77	40	(¥1)	ORIGINAL SOURCE: Synthetically Derived
				(x.)	PEATURE:
45 (2)		INFORMATION FOR SEQUENCE ID NO: 38:	A 7		(A) NAHE/KEY: Oligonucleotide Primer
	(£)	SEQUENCE CHARACTERISTICS:	į		
50		(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear	50		(B) LOCATION: Anneals to codons 222 to 228 of the DPA1 transcript of HLA Class II
	(11)	MOLECULE TYPE: Genomic DNA		(ix)	SEQUENCE DESCRIPTION: SEQ ID NO: 39:
Ę	(¥¥)	ANTI-SENSE: no	55		GGT CCC CTG GGC CCG GGG GTC 21

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(2) IMPORMATION FOR SEQUENCE ID NO: 41: (2) INFORMATION FOR SEQUENCE ID NO: 40: SEQUENCE CHARACTERISTICS: GCC AGA ACG CAG AGA CTT TAT SEQUENCE DESCRIPTION: SEQ ID NO: 40: FRAGMENT TYPE: Internal Fragment ANTI-SENSE: no MOLECULE TYPE: Genomic DNA (A) NAME/KEY: Oligonuclectide Primer DPA20 ANTI-SENSE: yes ORIGINAL SOURCE: FRAGMENT TYPE: Internal Fragment SEQUENCE CHARACTERISTICS: MOLECULE TYPE: Genomic DNA LENGTH: 21 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single
TOPOLOGY: Linear LOCATION: Anneals to codons 214 to 220 of the DPA1 transcript of HLA class II LENGTH: 21 base pairs
TYPE: Nucleic Acid
STRANDEDNESS: Single
TOPOLOGY: Linear Synthetically Derived 21 50 45 60 딿 30 25 15 5 u (2) INFORMATION FOR SEQUENCE ID NO: 42: Ê. ₹ (11) (Ť (¥¥) Œ Ē Ξ TTC TTG CAG CAG GAT AAG TA
Phe Leu Gln Gln Asp Lys Tyr
10 (B) LOCATION: Anneals to codons 7 to 13 of the DRB1 transcript of HLA class SEQUENCE DESCRIPTION: SEQ ID NO: 42: (A) NAME/KEY: Oligonaclectide Primer DRB23 FEATURE: ORIGINAL SOURCE: FRAGMENT TYPE: Internal Fragment SEQUENCE CHARACTERISTICS: (H ANTI-SENSE: no AAC TIG AAT ACC TIG ATC CAG Asn Leu Asn Thr Leu Ile Gln 70 MOLECULE TYPE: Genomic DNA SEQUENCE DESCRIPTION: SEQ ID NO: 41: (A) NAME/KEY: Oligonucleotide Primer DPA21 LENGTH: 20 base pairs TYPE: Nucleic Acid STRANDEDRESS: Single TOPOLOGY: Linear LOCATION: Anneals to codons 68 to 74 of the DPA1 transcript of HLA class II Synthetically Derived

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(A) NAME/KEY: Oligonucleotide Primer DRB25	ORIGINAL SOURCE: Synthetically Derived	FRACHENT TYPE: Internal Fragment	ANTI-SENSE: no	MOLECULE TYPE: Genemic DNA	(C) STRANDEDNESS: Single (D) TOPOLOGY: Linear	(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid	SEQUENCE CHARACTERISTICS:	INFORMATION FOR SEQUENCE ID NO: 44:		Fro Arg Fine Leu GLY Tyr Ser	CCA CGT TIC TING GAG TAC TCT 21	SEQUENCE DESCRIPTION: SEQ ID NO: 43:	of the DRB1 transcript of HLA class	(B) LOCATION: Anneals to codons 5 to 11	(A) NAME/KEY: Oligonucleotide Primer DRB24	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRACKENT TYPE: Internal Fragment	ANTI~SENSE: no	MOLECULE TYPE: Genomic DNA	DEDMI	(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid	SEQUENCE CHARACTERISTICS:	(2) INFORMATION FOR SEQUENCE ID NO: 43:	-79-	PCT/US92/01675
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(1)	INFORMATIO				(1 x)				(ix)	(¥1)	₹	(vì)	(11)			(±)	INFORMATI				(xi)					П
SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid	INFORMATION FOR SEQUENCE ID NO: 46:		30 35	AGA TGC ATC TAT AAC CAA GAG 21	SEQUENCE DESCRIPTION: SEQ ID NO: 45:	of the DRB1, DRB3, DRB4 and DRB5 transcripts of HLA class II	TOGRATON: Priests to codons	(A) NAME/KEY: Oligonucleotide Primer DRB16	FEATURE:	ORIGINAL SOURCE: Synthetically Derived	FRAGMENT TYPE: Internal Fragment	ANTI-SENSE: no	MOLECULE TYPE: Genomic DNA	(C) STRANDSUMESS: SINGLE (D) TOPOLOGY: Linear	(A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid	SEQUENCE CHARACTERISTICS:	(2) INFORMATION FOR SEQUENCE ID NO: 45:		10	TITC TIG GAG CAG GIT AAA CA 21 Arg Phe Leu Glu Gln Val Lys His	SEQUENCE DESCRIPTION: SEQ ID NO: 44:	II	(B) LOCATION: Anneals to codons 6 to 13	100-		PC1/US9Z/01675

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	į	(i x)	(41) (v)		(2) INFORMA (1)			(x1)		(*T)	(v1)	(v)	(1v)		
(B) LOCATION: Anneals to codons -8 to -2 of the DQB1 transcript of HLA class	(A) NAME/KEY: Oligonuclectide Primer DQB6		MOLECULE TYPE: Genomic DNA ANTI-SENSE: no FRAGMENT TYPE: Internal Fragment	(A) LENGTH: 21 base pairs (B) TYPE: Mucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear	INFORMATION FOR SEQUENCE ID NO: 47: (i) SEQUENCE CHARACTERISTICS:		AGA TAC TYC CAT AAC CAG GAG 21 Arg Tyr Phe His Asn Glu 35	SEQUENCE DESCRIPTION: SEQ ID NO: 46:	(B) LOCATION: Anneals to codons 29 to 35 of the DRB1, DRB3, DRB4 and DRB5 transcripts of HLA class II	(A) NAME/KEY: Oligonucleotide Primer	ORIGINAL SOURCE: Synthetically Derived	FRACMENT TYPE: Internal Fragment	ANTI-SENSE: no	(C) STRANDEDNESS: Single (D) TOPOLOGY: Linear	-81-
55	50	45	40	ы S	30		25	20	15		10		tn		
(ii) (iv)	(i)	(2) INFORMATI		(x1)		(1x)	(¥1)	(A)		(i)	(2) INFORMAT			(xt)	
TPE:	SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid	INFORMATION FOR SEQUENCE ID NO: 49:	CTG AGC TCC TCA CTG GCT GAG 21 Leu Ser Ser Ser Leu Ala Glu -5	(B) LOCATION: Anneals to codons -8 to -2 of the DQB1 transcript of HLA class II SEQUENCE DESCRIPTION: SEQ ID NO: 48:		FEATURE:	FRAGMENT TYPE: Internal Fragment ORIGINAL SOURCE: Synthetically Derived	AUTY-SENSE: no		QUENCE CHARACTERISTIC	INFORMATION FOR SEQUENCE ID NO: 48:		CTG AGC ACC CCA GTG GCT GAG 21 Leu Ser Thr Pro Val Ala Glu -5	D NO: 47	-822-1

- ₹ FRAGMENT TYPE: Internal Fragment
- (TV) ORIGINAL SOURCE: Synthetically Derived
- <u>*</u> FEATURE:
- (A) NAME/KEY: Oligonuclectide Primer DOB15

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- LOCATION: Anneals to codons -8 to -2 of the DQB1 transcript of HLA class
- È. SEQUENCE DESCRIPTION: SEQ ID NO: 49:

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CTG AGC ACC TCG GTG GCT GAG Lou Ser Thr Ser Val Ala Glu -5 21

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ÿ application, and the computer readable form of the same above "Sequence Listing" Section of the present submitted therewith, are the same. Applicants state that the paper copy of the

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WHAT IS CLAIMED:

- containing subject nucleic acid comprising: histocompatibility complex genotype of a subject in a sample A method for determining a major
- isolating nucleic acid from said sample;
- 9 amplifying said nucleic acid by polymerase oligonuclectide primer and at least one nonamplified with at least one conserved conserved oligonuclectide primer; gene locus and chromosome to be sequenced being chromosome to be sequenced being amplified with of said alleles for each gene locus and allele of said gene locus to be sequenced, all pair and at least one of said alleles for each polymerase chain reaction product for each chain reaction to generate sufficient at least one conserved oligonucleotide primer
- Î sequencing directly each polymerase chain reaction product for each allele at each gene that is sequenced; and and a conserved primer specific for each locus locus of each chromosome with Taq polymerase
- <u>a</u> analyzing each sequenced polymerase chain said subject. reaction product to determine the genotype of
- nucleic acid is genomic DNA. The method of claim 1 wherein said isolated
- prior to amplifying said nucleic acid: nucleic acid is RNA and further comprises the following step The method of claim 1 wherein said isolated
- synthesizing cDNA molecules for each allele of each gene locus to be sequenced, wherein said

conserved region of each allele of each said oligonucleotide primer that anneals to a synthesis employs a locus-specific

- histocompatibility genotype to be determined is a HLA Class II genotype. The method of claim 1 wherein said major
- gene locus to be sequenced is DQB1. The method of claim 4 wherein said Class II
- gene locus to be sequenced is DQA1. The method of claim 4 wherein said Class II
- gene loci to be sequenced are DRB 1/3/4/5. The method of claim 4 wherein said Class II
- gene loci to be sequenced is DPA 1. The method of claim 4 wherein said Class II
- gene loci to be sequenced is DPB 1. The method of claim 4 wherein said Class II
- each allele of such gene locus amplified with a conserved oligonuclectide primer pair to the nuclectide sequence of gene locus amplified with a conserved/non-conserved oligonucleotide primer pair. followed by comparing the sequence of each allele of each comparing the nucleotide sequence of each allele of each gene locus sequenced to known sequences for each such gene locus sequenced polymerase chain reaction product involves The method of claim 1 wherein analyzing said

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- Class II loci. nuclectide sequences of all allelss of all haplotypes for HLA conducted with a computer having a program including polymerase chain reaction product to determine genotype is The method of claim I wherein analyzing each
- said cDNA at about 37°C. includes annealing said conserved oligonuclectide primer to cDNA molecules with said conserved oligonuclectide primer 12. The method of claim 1 wherein said amplifying
- annealing said non-conserved primer to said cDNA at about cDNA molecules with said non-conserved primer includes 13. The method of claim I wherein said amplifying
- containing subject nucleic acid comprising: histocompatibility genotype of a subject in a sample 14. A method for determining the Class II
- (a) isolating nucleic acid from said sample;
- amplifying said nucleic acid by polymerase and at least one non-conserved oligonucleotide at least one conserved oligonuclectide primer chromosome to be sequenced being amplified with oligonucleotide primer pair and at least one of said alleles for each Class II gene locus and being amplified with at least one conserved II gene locus and chromosome to be sequenced allele of said Class II gene locus to be polymerase chain reaction product for each sequenced, all of said alleles for each Class chain reaction to generate sufficient
- <u>0</u> sequencing directly each polymerase chain reaction product for each allele at each Class

each Class II locus that is sequenced; and polymerase and a conserved primer specific for II gene locus of each chromosome with Tag

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- 3 determining the genotype of said subject by oligonuclectide primer. Class II locus amplified with a conserved nucleotide sequence of each allele of such degenerated oligonucleotide primer to the allele of each Class II locus amplified with a followed by comparing the sequence of each known sequences for each such Class II locus allele at each Class II locus sequenced to comparing the nuclectide sequence of each
- prior to amplifying said nucleic acid; nucleic acid is RNA and further comprises the following step The method of claim 14 wherein said isolated
- synthesizing cDNA molecules for each allele of oligonucleotide primer that anneals to a wherein said synthesis employs a locus-specific each Class II gene locus to be sequenced, Class II gene locus. conserved region of each allele of each said
- acid comprising: genotype of a subject in a sample containing subject nucleic 16. A method for determining the Class II HLA
- isolating total cellular RNA from said sample;
- synthesizing cDNA molecules for each allele of each said Class II gene locus; anneals to a conserved region of each allele of sequenced, wherein said synthesis employs a locus-specific oligonucleotide primer that at least one Class II gene locus to be

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- amplifying said cDNA molecules by polymerase sequenced being amplified with at least one Class II gene locus and chromosome to be pair and at least one of said alleles for each at least one conserved oligonucleotide primer chromosome to be sequenced being amplified with alleles for each Class II gene locus and II gene locus to be sequenced, all of said reaction product for each allele of said Class chain reaction to generate a polymerase chain
- <u>@</u> sequencing directly each polymerase chain allele; and produce a nucleic acid sequence ladder for each each Class II locus that is sequenced to polymerase and a conserved primer specific for II gene locus of each chromosome with Tag reaction product for each allele at each Class

one non-conserved oligonucleotide primer; conserved oligonucleotide primer and at least.

- analyzing each nucleic acid ladder to determine sequence of each allele of such Class II locus oligonucleotide primer pair to the nucleotide locus amplified with a conserved/non-conserved each such Class II locus followed by comparing the sequence of each allele of each Class II Class II locus sequenced to known sequences for nucleotide sequence of each allele of each the genotype of said subject by comparing the amplified with a conserved oligonucleotide
- genes of said subject. for the DRB1, DRB3, DRB4, DRB5, DQB1, DQA1, DPA1 and DPB1 HLA genotype to be determined includes nucleotide sequences 17. The method of claim 16 wherein said Class II

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- transcript. strand of DNA which anneals to codons 105 to 111 of the DQB 18. An oligonucleotide primer comprising a single
- GGTGGTTGAGGGCCYCTGTCC. (SEQ. ID NO:1) 19. An oligonuclectide primer having the sequence
- strand of DNA which anneals to codons 1 to 7 of DQB 20. An oligonucleotide primer comprising a single
- AGAGACTCTCCCGAGGATTTC. 21. An oligonucleotide primer having the sequence (SEQ. ID NO:4)
- strand of DNA which anneals to codons 148 to 155 of its DQA 22. An oligonuclectide primer comprising a single
- GGTGAGGTTACTGATCTTGAAG. (SEQ. ID NO:3) 23. An oligonuclectide primer having the sequence
- strand of DNA which anneals to codons -10 to -4 of DQA cDNAs. 24. An oligonucleotide primer comprising a single
- CTGTCCTCCGTGATGAGCCC. (SEQ. ID NO:7) An oligonuclectide primer having the sequence
- transcript. strand of DNA which anneals to codons 105 to 111 of DRB1 An oligonuclectide primer comprising a single
- GTGCTGCAGGGGCTGGGTCTT. An oligonucleotide primer having the sequence (SEQ. ID NO:2)

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- strand of DNA which anneals to codons -33 to -26 of DRB1 transcripts. 28. An oligonucleotide primer comprising a single
- TGTTCTCCAGCATGGTGTGTC. An oligonucleotide primer having the sequence (SEQ. ID NO:6)
- strand of DNA which anneals to codons 7 to 13 of DRB1 transcript. An oligonucleotide primer comprising a single
- TTCTTGCAGCAGGATAAGTA. (SEQ. ID NO:42) An oligonucleotide primer having the sequence
- transcript. strand of DNA which anneals toe codons 5 to 11 of DRB 1 32. An oligonuclectide primer comprising a single
- CCACGTTTCTTGGAGTACTCT. An oligonucleotide primer having the sequence (SEQ. ID NO:43)
- transcript. strand of DNA which anneals to codons 6 to 13 of DRB 1 34. An oligonuclectide primer comprising a single
- TTTCTTGGAGCAGGTTAAACA. An oligonuclectide primer having the sequence (SEQ. ID NO:44)
- transcript. strand of DNA which anneals to codons 105 to 111 of DPB An oligonuclectide primer comprising a single
- GITGEGGECTGCAAGGGCCC. 37. An oligonucleotide primer having the sequence (SEQ. ID NO:24)

- 38. An oligonuclectide primer comprising a single strand of DNA which anneals to codons 222 to 228 of DPA transcript.
- 39. An oligonucleotide primer having the sequence GGTCCCCTGGGCCCCGGGGGTC. (SEQ. ID NO:39)
- 40. An oligonuclectide primer comprising a single strand of DNA which anneals to codons -19 to -13 of DPB transcript.
- 41. An oligonucleotide primer having the sequence CGGACAGTRGCTCTGACGGCG. (SEQ. ID NO:23)
- 42. An oligonuclectide primer comprising a single strand of DNA which anneals to codons -23 to -17 of DPA transcript.
- 43. An oligonucleotide primer sequence of CATATCAGAGCTGTGATCTTG. (SEQ. ID NO.32)
- 44. An oligonucleotide primer comprising a single strand of DNA which anneals to codons 59 to 65 of DPA transcript.
- 45. An oligonucleotide primer having the sequence CTGGCTAACAFTGCTATATTG. (SEQ. ID NO:38)
- 46. An oligonuclectide primer comprising a single strand of DNA which anneals to codons 104 to 110 of DPA transcript.
- 47. An oligonucleotide primer having the sequence GTCAATGTGGCAGATGAGGGT. (SEQ. ID NO.31)

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- 48. An oligonuclectide primer having the sequence GCCGCTGCACTGTGAAGCTC. (SBQ. ID NO:13)
- 49. An oligonucleotide primer having the sequence CCTGGAGGGGGAAACATTCAC. (SEQ. ID NO:11)
- 50. An oligonucleotide primer having the sequence gracergrerecacac. (SEQ. ID NO:12)
- 51. An oligonuclectide primer having the sequence CACGGTTCCGGTAGCAGCGGTAG. (SEQ. ID NO:14)
- 52. An oligonuclectide primer having the sequence CCTGGAGGGGGAAACATTCAC. (SEQ. ID NO:25)
- 53. An oligonucleotide primer having the sequence CTTGGGAAACACGGTCACCTC. (SEQ. ID NO:33)
- 54. An oligonuclectide primer having the sequence GCCAGAACGCAGACTTTAT. (SEQ. ID NO:40)
- 55. An oligonucleotide primer comprising a single strand of DNA which anneals to base pairs 18 to 38 of intron 3 of DRB loci.
- 56. An oligonuclectide primer having the sequence of GCCAAGAGTGGGGCTCGCAGC. (SEQ. ID NO:20)
- 57. An oligonucleotide primer comprising a single strand of DNA which anneals to codons -4 to +3 of the DRB transcript.
- 58. An oligonucleotide primer having the sequence CTGGCTTTGGCTGGGGACACC. (SEQ. ID NO:5)

- strand of DNA which anneals to codons 88 to 94 of the DQB 59. An oligonuclectide primer comprising a single
- TOGCOTOTGCAGGGTCGCGCG. 60. An oligonuclectide primer having the sequence (SEQ. ID NO:8)
- strand of DNA which anneals to codons 11 to 17 of the DQB transcript. An oligonuclectide primer comprising a single
- of TTTAAGGGCATGTGCTACTTC. 62. An oligonucleotide primer having the sequence (SEQ. ID NO:9)
- intron 2 of the DPB locus. strand of DNA which anneals to base pairs -42 to -62 of 63. An oligonuclectide primer comprising an single
- AGAGGGAGAAAGAGGATTAGA. (SEQ. ID NO:27) 64. An oligonucleotide primer having the sequence
- strand of DNA which anneals to intron 39 to 59 of intron 3 of the DPB gene. An oligonuclectide primer comprising a single
- GCCCTGGGCACGGGCCCGCGG. (SEQ. ID NO:28) 66. An oligonucleotide primer having the sequence
- strand of DNA which anneals to base pairs -69 to -50 of intron 2 of the DPA1 locus. 67. An oligonuclectide primer comprising a single
- CTCTAGCTTTGACCACTTGC. 68. An oligonuclectide primer having the sequence (SEQ. ID NO:35)

- 3 of the DPA1 locus. strand of DNA which anneals to base pairs 55 to 71 of intron 69. An oligonuclectide primer comprising a single
- AGTCTGAGGGTGGCAGAGAGG. 70. An oligonuclectide primer having the sequence (SEQ. ID NO:36)
- transcript. strand of DNA which anneals to codons 87 to 94 of the DRB An oligonucleotide primer comprising a single
- strand of DNA which anneals to codons 38 to 45 of the DRB 72. An oligonucleotide primer comprising a single
- GCGCTTCGACAGCGACGTGG. An oligonuclectide primer having the sequence (SEQ. ID NO:16)
- CGGCCCAAAGCCCTCACTCAC. An oligonucleotide primer having the sequence (SEQ. ID NO:29)
- 75. An oligonucleotide primer having the sequence GGCCTGAGTGTGGBAGG. (SEQ. ID NO:37)
- TACTGATGGTGCTCACAT. (SEQ. ID NO:26) 76. An oligonuclectide primer having the sequence
- CGCTCATGTCCGCCCCCCCCC. (SEQ. ID NO:30) 77. An oligonucleotide primer having the sequence
- CTGCTGAGTCTCCGAGGAGCT. (SEQ. ID NO:34) An oligonucleotide primer having the sequence
- AACTIGAATACCTTGATCCAG. 79. An oligonucleotide primer having the sequence (SEQ. ID NO:41)

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ATGGGGAGATGGTCACTGTGG. (SEQ. ID NO:10) 80. An oligonucleotide primer having the sequence

- TACGGTCCCTCTGGCCAG. (SEQ. ID NO:15) An oligonuclectide primer having the sequence
- in a sample containing subject nucleic acid comprising: major histocompatibility complex class genotype of a subject 82. A method for rapid automated determination of
- (a) isolating nucleic acid from said sample with an RNA/DNA extractor;
- 9 amplifying said nucleic acid by polymerase oligonuclectide primer; oligonucleotide primer and one non-conserved gene locus and chromosome to be sequenced being pair and at least one of said alleles for each at least one conserved oligonucleotide primer chromosome to be sequenced being amplified with of said alleles for each gene locus and chain reaction using a thermocycler to generate amplified with at least one conserved allele of each gene locus to be sequenced, all a polymerase chain reaction product for each
- <u>@</u> sequencing directly each polymerase chair reaction product for each allele at each gene conserved primer specific for each locus to be sequencing apparatus with Tag polymerase and a sequenced; and locus of each chromosome in an automated
- a) analyzing each sequenced polymerase chain the sequence of each allele of each gene locus with allelic sequence information to compare said subject with a computer having a data base reaction product to determine the genotype of

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pair to the nucleotide sequence of each allele oligonucleotide primer pair. conserved/non-conserved oligonucleotide primer of such gene locus amplified with a conserved each allele of each gene locus amplified with a locus followed by comparing the sequence of sequenced to known sequences for each such gene

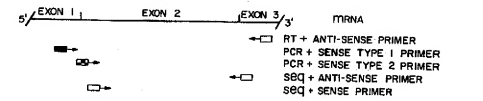
- more polymorphic gene locus of a subject in a sample containing subject nucleic acid comprising: A method for determining the genotype at one or
- isolating nucleic acid from said sample;
- ਉ amplifying said nucleic acid by polymerase oligonuclectide primer and at least one nongene locus and chromosome to be sequenced being pair and at least one of said alleles for each allele of said gene locus to be sequenced, all conserved oligonucleotide primer; amplified with at least one conserved at least one conserved oligonuclectide primer chromosome to be sequenced being amplified with of said alleles for each gene locus and polymerase chain reaction product for each chain reaction to generate sufficient
- 9 sequencing directly each polymerase chain reaction product for each allele at each gene enzyme and a conserved primer specific for each locus that is sequenced; and locus of each chromosome with a sequencing
- <u>a</u> analyzing each sequenced polymerase chain reaction product to determine the genotype of said subject.
- nucleic acid is genomic DNA. The method of claim 63 wherein said isolated

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- prior to amplifying said nucleic acid: nucleic acid is RNA and further comprises the following step 85. The method of claim 83 wherein said isolated
- synthesizing cDNA molecules for each allele of conserved region of each allele of each said oligonuclectide primer that anneals to a synthesis employs a locus-apecific gene locus. each gene locus to be sequenced, wherein said
- subject in a sample containing subject nucleic acid comprising: the genotype at one or more polymorphic gene locus of a 86. A method for rapid automated determination of
- (a) isolating nucleic acid from said sample with an RNA/DNA extractor;
- oligonuclectide primer; amplifying said nucleic acid by polymerase oligonuclectide primer and one non-conserved gene locus and chromosoms to be sequenced being pair and at least one of said alleles for each of said alleles for each gene locus and amplified with at least one conserved at least one conserved oligonucleotide primer chromosome to be sequenced being amplified with allels of each gene locus to be sequenced, all a polymerase chain reaction product for each chain reaction using a thermocycler to generate
- <u>0</u> sequencing directly each polymerase chain sequencing apparatus with a sequencing enzyme reaction product for each allele at each gene to be sequenced; and and a conserved primer specific for each locus locus of each chromosome in an automated

Ē analyzing each sequenced polymerase chain conserved/non-conserved oligonucleotide primer oligonucleotide primer pair. of such gene locus amplified with a conserved pair to the nucleotide sequence of each allele each allele of each gene locus amplified with a sequenced to known sequences for each such gene the sequence of each allele of each gene locus with allelic sequence information to compare said subject with a computer having a data base reaction product to determine the genotype of locus followed by comparing the sequence of

FIG. IA



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FIG. IB

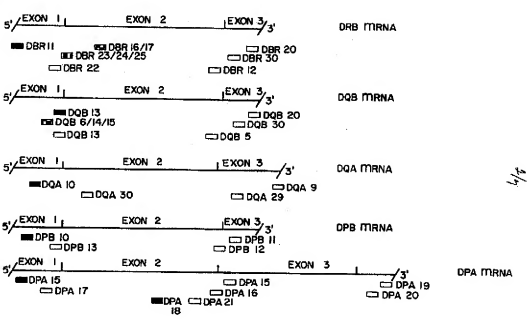
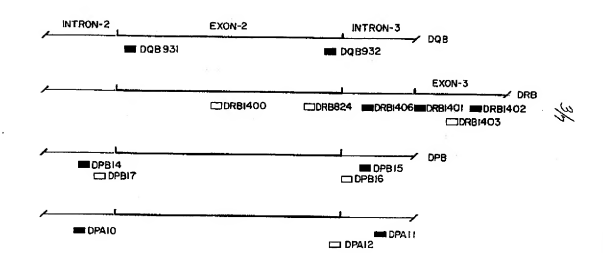
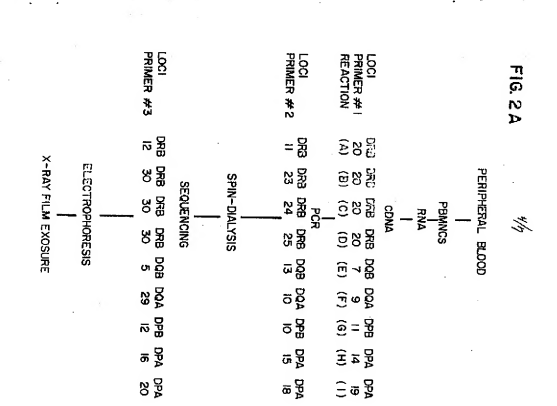


FIG. I C





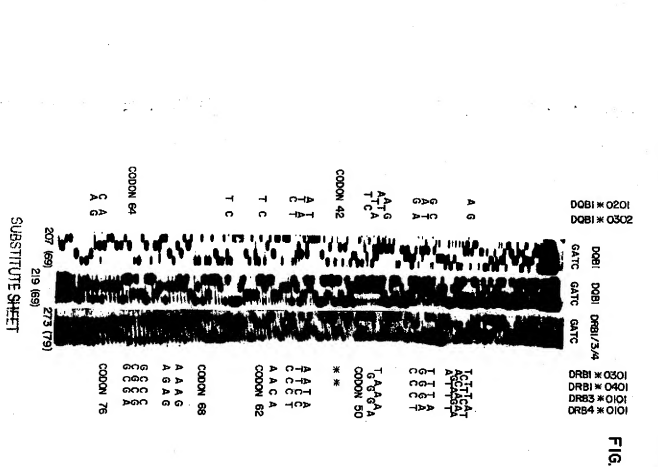
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FIG. 2B

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FORENSIC SAMPLES

DNA



LOCI PRIMER #3

DRB DRB DRB DRB DQB DPB 12 1400 1400 1400 5 16

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SEQUENCING

LOCI
PRIMER #1
PRIMER #2
REACTION

DRB DRB DRB DRB DQB I I406 I406 I406 I406 932 22 23 23 24 93I (S) (V) (T) (U) (W)

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3=08

SPIN-DIALYSIS

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X-RAY FILM EXPOSURE

ELECTROPHORESIS

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FIG. 4

INTERNATIONAL SEARCH REPORT

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	Y CLINICAL CHEMISTRY, 1989, I.J. MCBride Methods Involving 2196-2201, 308 espe-	Y PROCEEDINGS OF THE Volume 85, issued a "Direct Sequencing Genomic DNA", pages	Y Description of al., pages 141-144,	y BIOTECHNIQUES GOTMAN et al Amplification 325-329, see	Catagory* Citation of Document	III. DOCUMENTS CONSIDERED TO BE RELEVANT "		U.S.	Classification System	Mark Superior Contract	According to intermedical Parent Classification [Fre [5]: C120 1/68; C122 19/14 US CL : 415/6, 91
Special component of cited decuments: It is an experiment published effect the interruptional filling document defining the greener state of the art which is not considered to be of periodic relevance and the process of the art which is application but of led to an advant the principle on the process of t	HEMISIRY, Volume 38 Mo. 11, issued Movember McGride et al., "Automated DNA Sequencing prolying Polymerase Chain Reaction", pages see especially pages 2200-2201.	DINGS OF THE MATIONAL ACADEMY OF SCIENCE USA, 85, issued January 1988, D.R. Engelke et al., t Sequencing of Enzymatically Amplified Human C DNA", pages 544-548, see entire document.), Volume 31, issued March 1990, S.G.B. "HLA-Dhb Mucleotide Sequences, 1990", see entire document.	BIOTECHNIQUES, Volume 7, No. 4, issued April 1989, K.B. Gorman et al., 'Simplified Method for Selective Amplification and Direct Sequencing of cDNAs' pages 326-329, see entire document.	Citation of Document, 16 with indication, where appropriate, of the relevant passages 17	D TO BE RELEVANT 14	Documentation Searched other than Minkman Documentation to the extent that such Documents are included in the Fields Searched		Classification Symbols	Minimum Depurpentation Searched	According to international Privit Classification (IPC) or to both National Classification and IPC ISC [5]: C120 1/6; C129 19/14 US CL: + 435/6; 9:
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